

# Online Research @ Cardiff

This is an Open Access document downloaded from ORCA, Cardiff University's institutional repository: <https://orca.cardiff.ac.uk/id/eprint/122803/>

This is the author's version of a work that was submitted to / accepted for publication.

Citation for final published version:

Wild, John ORCID: <https://orcid.org/0000-0003-3019-3889>, Smith, Phil E. M. ORCID: <https://orcid.org/0000-0003-4250-2562> and Knupp, Carlo ORCID: <https://orcid.org/0000-0001-9127-2252> 2019. Objective derivation of the morphology and staging of visual field loss associated with long-term vigabatrin therapy. *CNS Drugs* 33 (8) , pp. 817-829. 10.1007/s40263-019-00634-2 file

Publishers page: <https://doi.org/10.1007/s40263-019-00634-2>  
<<https://doi.org/10.1007/s40263-019-00634-2>>

Please note:

Changes made as a result of publishing processes such as copy-editing, formatting and page numbers may not be reflected in this version. For the definitive version of this publication, please refer to the published source. You are advised to consult the publisher's version if you wish to cite this paper.

This version is being made available in accordance with publisher policies.

See

<http://orca.cf.ac.uk/policies.html> for usage policies. Copyright and moral rights for publications made available in ORCA are retained by the copyright holders.



# CNS Drugs

## Objective derivation of the morphology and staging of visual field loss associated with long-term vigabatrin therapy --Manuscript Draft--

<b>Manuscript Number:</b>	CNSA-D-19-00041R1
<b>Full Title:</b>	Objective derivation of the morphology and staging of visual field loss associated with long-term vigabatrin therapy
<b>Article Type:</b>	Original Research Article
<b>Funding Information:</b>	
<b>Abstract:</b>	<p><b>Background</b> The morphology and between-eye symmetry of the visual field loss associated with the anti-epileptic drug vigabatrin (VAVFL) has received little attention. <b>Objective</b> To model the appearance, and ensuing staging, of VAVFL, derived with the European Medicines Agency approved perimetric protocol.</p> <p><b>Methods</b> A retrospective, cross-sectional, observational design identified 123 adults who had received vigabatrin for refractory seizures and who had no evidence of co-existing retino-geniculo-cortical visual pathway abnormality. Thirty-eight adults with refractory seizures and identical inclusion criteria, but no exposure to vigabatrin, acted as controls. For each group, the median outcome at each stimulus location in each eye (of absolute loss, relative loss or Pattern Deviation probability level, as appropriate) was derived for each successive ten pairs of fields, ranked for severity. Between-eye symmetry was quantified by an index which accounted for severity of loss and which was referenced to the likelihood of the occurrence of symmetry due to chance.</p> <p><b>Results</b> The modelled VAVFL was bilateral and highly symmetrical and was described by six stages which were all independent of the extent of vigabatrin exposure. The loss originated in the extreme temporal periphery and encroached centripetally along all meridians towards fixation. The initial appearance within the central field (Stage Two) occurred inferior-nasally. Subsequent stages exhibited increasing loss which was greater nasally than temporally. Stage Six described concentric loss extending to approximately 15° eccentricity from fixation.</p> <p><b>Conclusion</b> The model exhibited a consistent pattern of VAVFL. The staging of the loss could assist the risk:benefit analysis of vigabatrin for the treatment of epilepsy.</p>
<b>Corresponding Author:</b>	John Wild, PhD, DSc Cardiff University Cardiff, UNITED KINGDOM
<b>Corresponding Author Secondary Information:</b>	
<b>Corresponding Author's Institution:</b>	Cardiff University
<b>Corresponding Author's Secondary Institution:</b>	
<b>First Author:</b>	John Wild, PhD, DSc
<b>First Author Secondary Information:</b>	
<b>Order of Authors:</b>	John Wild, PhD, DSc Phillip E M Smith, MD, FRCP Carlo Knupp, PhD
<b>Order of Authors Secondary Information:</b>	
<b>Author Comments:</b>	<p>Given the issue with one of the referees of our last manuscript, we would respectfully insist that the manuscript is NOT refereed by anyone who has previously declared a conflict of interest with Lundbeck LLC and/ or with Ovation Pharmaceuticals, Inc.</p> <p>The referees who reviewed the 'appealed' version of our last manuscript were both interested in, and knowledgeable about, vigabatrin toxicity. Perhaps they could be</p>

	<p>approached again?</p>
<b>Response to Reviewers:</b>	<p>CNSA-D-19-00041</p> <p>Objective derivation of the morphology and staging of visual field loss associated with long-term vigabatrin therapy</p> <p>We acknowledge the contribution from each of the reviewers. The review process has resulted in a much better manuscript.</p> <p>All changes included in the revised manuscript are highlighted in green and are described below.</p> <p>Response to Reviewer #1</p> <p>1.The subjects are adults. The authors have avoided discussing children but do they have any sense of whether this data would be of alarm to younger age groups. We did not 'avoid(ed) discussing children'. The case series was compiled from adults on the basis of their ability to perform perimetry.</p> <p>We have inserted a paragraph in the Discussion at the bottom of page 14/ top of page 15 of the revised manuscript.</p> <p>The model was developed from the visual fields of adults. The reduction in amplitude of the 30Hz flicker cone electroretinogram (ERG) in infants treated with vigabatrin for infantile spasms [31] is compatible with that for adults manifesting VAVFL [32]. Similarly, the topographical characteristics of the reduced peripapillary retinal nerve fibre layer thickness in children [33] is also compatible with that found in adults [34-38]. Those with vigabatrin-associated 30Hz flicker cone ERG abnormality in infancy subsequently manifest VAVFL, and corresponding retinal nerve fibre layer thinning, in later childhood/ early adolescence [39]. There is no reason to suggest, therefore, that the vigabatrin toxicity manifested in infancy will result in a different appearance to the VAVFL when the latter measure is obtained in later life.</p> <p>2.The reader would like to know, just generally, how impaired these subjects were? Were some in wheelchairs? Head control or eye movement control problem? Any with frequent seizures e.g., more than a few per day.</p> <p>The Reviewer seems to have overlooked our commentary at the top of page 17 of the initial version of the manuscript, namely:</p> <p>All 14 individuals exhibited impaired mobility and reported symptoms explainable by their field loss, the most common of which was bumping into individuals in crowded locations.</p> <p>Nevertheless, we have modified the paragraph (now top of page 19 of the revised manuscript). The new text is given in non-italicised text.</p> <p>All 14 individuals exhibited profoundly impaired mobility and reported symptoms explainable by their VAVFL, the most common of which was bumping into individuals in crowded locations. However, some individuals with Stage 5 VAVFL also reported a negative impact on their activities of daily living.</p> <p>We have also inserted a sentence in the Methodology at the end of the paragraph on Page 7</p> <p>All individuals had been seizure free for a minimum of 24 hours prior to the visual field examination.</p> <p>3.I don't see any data on false-positive and negatives to the field tests – were there any? This makes it easier to judge the validity of the tests.</p>

We made reference to the criteria for incorrect responses to the false-positive and false-negative catch trials the field tests in the initial version of the manuscript (page 7, middle paragraph).

The reliability criteria comprised  $\leq 20\%$  incorrect responses to the fixation loss catch trials and  $\leq 15\%$  incorrect responses to the false-positive catch trials. A criterion of  $\leq 30\%$  incorrect responses was used for the false-negative catch trials although this was widened with increasing severity of the field loss [25].

Accordingly, we have not revised the manuscript in this regard.

4. Do the authors have any data on visible changes to retina morphology that accompanied the field loss. And if so what stage did these develop.

We have inserted the following in the Discussion from the middle of page 15 onwards

The most common bilateral finding by fundoscopy, through a dilated pupil, was either generalised or localised arteriolar narrowing which was noted at all stages of VAVFL. However, this finding was occasionally found in the presence of a normal field and, therefore, may well be associated with vigabatrin usage. Subtle bilateral retinal nerve fibre layer changes and optic nerve head pallor were noted from Stage 4 onwards. A variety of bilateral peripheral degenerative changes were also noted including hypopigmented/ white spots and surface wrinkling. However, there was considerable variation in the fundal appearance between individuals within a given stage. The various features are in agreement with those reported previously [3, 22, 40-41] were often subtle and suggest that there is a wide spectrum of potential retinopathy associated with vigabatrin toxicity. There was no evidence of these findings in the individuals with no exposure to vigabatrin. Twenty-nine individuals exposed to vigabatrin had undergone optical coherence tomography of the peripapillary retinal nerve fibre layer at the time of the examination. All three individuals who manifested a normal visual field exhibited a normal nerve fibre layer thickness. Of the three individuals who had Stage 1 loss, two exhibited a normal thickness. An abnormally thin nerve fibre layer, characteristic of vigabatrin toxicity [34-38], was present from Stage 2 onwards in 22 individuals. The one remaining individual manifested early Stage 2 VAVFL and a normal a normal nerve fibre layer thickness.

5. Were there any other reasons for exclusion...

The exclusion criteria are listed in the methods; no other exclusion criteria were applied post selection.

6. Do the authors have any sense of what happens if vigabatrin is stopped. Do field changes continue to progress

Please see the response to Query #1 from Reviewer #2. The following text has been inserted into the manuscript.

An unpublished audit of the long-term follow-up, over a maximum of eight years, of individuals in the current study indicates that the field loss neither improves nor deteriorates following withdrawal of vigabatrin.

Minor change

We have incorporated 'No' into the abstract.

Response to Reviewer #2

Major Comments

1. The major weakness of this study is failure to provide data in patients followed up longitudinally over time. Although the authors imply that patients progress from Stage 1

(least severe) to Stage 6 (most severe), they have not confirmed that such progression occurs. Most importantly, how can staging data be interpreted without knowledge of the probability of progression at each stage, and an estimate of the rate of progression? These concerns are reinforced by the finding that the occurrence of visual field defects was related to cumulative dose and duration of treatment, but there was no correlation between stages and exposure. The authors' comment that toxicity is 'idiosyncratic' does not explain the lack of correlation between severity of the defect and dose/duration of exposure. These issues need to be discussed.

We have inserted an essentially new paragraph in the Discussion on Page 17. The new text is given in non-italicised text.

The staging of VAVFL was derived from cross-sectional evidence and does not imply progressive loss. Due to the potency of the potential toxicity, most individuals had been withdrawn from vigabatrin either immediately prior to the introduction of the visual field examination into the care regime or following confirmation of the VAVFL. As such, neither the probability of progression at each stage nor an estimate of the rate of progression can be determined. However, the presumption is that, given continued vigabatrin therapy, VAVFL will progress through the various stages. The lack of an association between the stage of loss, at detection, and either the duration or the cumulative dose of vigabatrin implies that the relationship between the extent of exposure and both the onset and the severity of VAVFL varies depending upon the individual susceptibility to vigabatrin. However, the time of detection is not the time to onset of the VAVFL. The rate of any subsequent progression, therefore, remains unknown. A case of progressive loss during approximately 7.75 years of vigabatrin therapy and illustrated in terms of the Pattern Deviation probability map of the Central 24-2 Threshold Test and, subsequently, the C30-2T is shown in Fig 6. The outcome of the corresponding FF135 at the final visit is shown in Online Resource Fig 4. VAVFL has also been shown to be progressive whilst on therapy in all 14 cases over a mean follow-up of 10.7 years with the rate of reduction in the I4e isopter of kinetic perimetry increasing with increase in cumulative dose [6]. An unpublished audit of the long-term follow-up, over a maximum of eight years, of individuals in the current study indicates that VAVFL neither improves nor deteriorates following withdrawal of vigabatrin.

## 2. Selection bias

We have inserted a sentence relating to selection bias in the last sentence on Page 6.

There was no conscious selection bias. Patients were included on the basis of sequential retrieval of cases which met the inclusion/ exclusion criteria.

## 3. Continuity in severity of field loss across stages. How were cutoffs across stages (and the actual number of stage) defined?

We have modified the paragraph (now at the bottom of page 9/ top of page 10 of the revised manuscript). The new text is given in non-italicised text.

The stages were empirically selected on the basis of clinically meaningful intervals of peripheral field loss and, subsequently, of central field loss. These intervals, in turn, were based upon the magnitude of the between-examination physiological variability in the differential light sensitivity as a function of severity of loss [28-29]. The derivation ensured continuity across stages.

## 4. Please clarify whether the ophthalmologist was aware of the exposure status.

We have inserted a commentary on page 15.

The ophthalmological examinations were undertaken by any one of four ophthalmologists, depending upon the particular clinic, who were all unaware of the findings from their colleagues. All four were aware of the anti-epileptic drug history and, usually, of the visual field. However, the modelling of the visual fields from those exposed to vigabatrin and from the control individuals was objective and independent of the outcome from the ophthalmological and neurological examinations.



#### Compile Table about characteristics/ Statistical differences

We have inserted the summary statistics for the duration and cumulative dose of vigabatrin into the penultimate paragraph on page 12 of the revised manuscript.

Those with VAVFL manifested a greater cumulative dose (mean 7.94kg; SD 4.45; range 1.1 to 20.7) and a longer duration of therapy (8.86 years; SD 3.51; range 0.66 to 16.05) at the time of perimetry than those without field loss (mean 3.36kg; SD 4.77; range 0.11 to 16.14; and mean 3.62 years; SD 3.94; range 0.33 to 11.83; respectively). The difference between means (Student's t-test for two independent samples) were 4.58 kg (95% CI 1.71-7.45;  $p < 0.001$ ) and 5.25 years (95% CI 2.98-7.53;  $p < 0.001$ ), respectively.

We have inserted and additional sentence on Page 12.

There was no association between the stage of VAVFL and either age, age at onset of epilepsy or age at onset of vigabatrin.

#### Minor Comments

##### 1.Remove without prescription

The statement concerning the free availability on-line without prescription has been removed from the revised manuscript

2.Lines 26-50 This paragraph does not describe appropriately the background for the study.

The Reviewer is incorrect in his/ her commentary.

i)The best description of the methodology used to compile the Registry described by Krauss et al (2016) is that of Pellock et al (2011) [Epilepsy & Behavior 2011;22:710-717].

Page 2 Paragraph 4 of Pellock et al (2011) states that:

'Regular ophthalmologic assessments are required throughout vigabatrin therapy: at baseline ( $\leq 4$  weeks after therapy initiation), at least every 3 months during therapy and 3 to 6 months after discontinuation. Fulfilment of this mandate is documented by registry receipt of an ophthalmologic assessment form.....In many instances, the first actual vision test may be conducted well after entry into the registry.'

The Registry was an FDA mandated requirement for the marketing authorisation of Sabril within the USA. The data analysis for the Krauss et al (2016) paper was undertaken by a third party institution and the editorial support for the paper was funded by the Marketing Authorisation Holder for Sabril in the USA. The paper is carefully worded, is ambiguous in crucial places, and largely concentrates on the number of individuals enrolled within the Registry and the likelihood of retention of these individuals over time. The referee is correct in his/ her assertion that the paper by Krauss et al (2016) 'does not provide (sic) baseline data on visual fields'; however, the paper states that:

'Clinicians reported that eight patients discontinued treatment because of visual field defects (VFD). The patients ranged from ages 18 to 55 years at entry into the SHARE program and were divided equally by gender. One 19 year old patient was reported to have an indication of IS, the other patients had CPS. Three patients entered the program with previous vigabatrin exposure. The five patients who were newly treated had vigabatrin discontinued by their clinicians because of VFD after exposures of 13 months to 3.3 years (Mean: 23 months).'

T

he clear implication from the above is that the visual field loss in at least five patients occurred after the baseline examination.

There is no discussion in the paper on the (low) prevalence of visual field loss and no

	<p>mention of the fact that 'the data are not interpreted as indicating lack of visual toxicity.' Although not stated, the clear implication to the reader is that the prevalence of the toxicity, derived from a [this] prospective study, is much lower than that derived from the (European) retrospective studies.</p> <p>ii)The Sergott et al (2016) paper is described as a 'small study sponsored by Lundbeck.' It should be noted that this study was a one year prospective study was mandated by the FDA.</p> <p>Given the above and given the acceptance by Reviewer #1, we have not altered the paragraph.</p> <p>Response to Comments from Editor</p> <ol style="list-style-type: none"> <li>1.We have moved the definition of VAVFL to the end of the figure legend</li> <li>2.We have renumbered the figures as advised</li> <li>3.We have revised the format of the supplementary content as advised</li> </ol> <p>Our preference would be to embed a link to the videos within the article.</p>
<b>Suggested Reviewers:</b>	<p>Mark Lawden, PhD, FRCP Consultant Neurologist, University Hospitals Of Leicester NHS Trust mark.lawden@uhl-tr.nhs.co.uk International authority on vigabatrin toxicity</p> <p>Emilio Perruca, MD, PhD Professor. Director, University of Pavia. IRCCS. Mondino National Neurological Institute in Pavia perucca@unipv.it International reputation for the neurology and clinical pharmacology of epilepsy</p> <p>James Acheson, FRCOPhth, FRCP National Hospital for Neurology and Neurosurgery and Moorfields Eye Hospital, London james.acheson@nhs.net Internationally renown clinical neuro-ophthalmologist with an interest in vigabatrin toxicity</p> <p>Lars Frisen, MD Institute of Neuroscience and Physiology, University of Gothenburg lars.frisen@neuro.gu.se Internationally acclaimed neuro-ophthalmologist with a knowledge of vigabatrin toxicity</p> <p>Ivan Goldberg, MD Professor, University of Sydney ivan.goldberg@sydney.edu.au Internationally acclaimed ophthalmologist with a special interest in perimetry</p>



**Cardiff Centre For Vision Sciences Research  
College of Biomedical and Life Science**

**Cardiff University**

Maindy Road  
Cardiff CF24 4HQ  
Wales, UK

Tel +44(0)29 2087 4374  
Fax +44(0)29 2087 4358

[www.cardiff.ac.uk](http://www.cardiff.ac.uk)

Sue Pochon  
Editor  
CNS drugs

22nd February 2019

Dear Sue

**Prifysgol Caerdydd**

Heol Maindy  
Caerdydd CF24 4HQ  
Cymru, Y Deyrnas Unedig

Ffôn +44(0)29 2087 4374  
Ffacs +44(0)29 2087 4358

[www.caerdydd.ac.uk](http://www.caerdydd.ac.uk)

As promised in an earlier email relating to our last manuscript on vigabatrin (which is now published in *CNS Drugs*), please find uploaded a new manuscript entitled:

**Objective derivation of the morphology and staging of visual field loss associated with long-term vigabatrin therapy**

which we hope will be found suitable for publication in *CNS Drugs*.

The manuscript is particularly aimed at neurologists, neuro-ophthalmologists, general ophthalmologists and other healthcare workers associated with the care of patients with epilepsy.

Due to the cross-discipline nature of the topic, and given our past publications in the journal, we feel that *CNS Drugs* is the most appropriate setting for the above readership.

As you know vigabatrin is a highly effective drug for the treatment of refractory focal seizures but, for almost two decades, has been associated with visual field loss and with various other retinal/optic nerve structural and functional abnormalities.

Inexplicably, the appearance of vigabatrin-associated visual field loss (VAVFL) has never been illustrated/ described in detail.

The manuscript is topical in that:

- The use of vigabatrin is likely to increase, given



Registered Charity, no. 1136855  
Elusen Gofrestredig, rhif 1136855



- the apparent lack of evidence for the toxicity arising from the two recent prospective studies (but these only involved *short-term* exposures)
- the drug has recently become available as a generic
- the drug is also freely obtainable on-line without prescription

**Prifysgol Caerdydd**

Heol Maindy  
Caerdydd CF24 4HQ  
Cymru, Y Deyrnas Unedig

Ffôn +44(0)29 2087 4374  
Ffacs +44(0)29 2087 4358

[www.caerdydd.ac.uk](http://www.caerdydd.ac.uk)

The manuscript is novel in that it:

- Presents the first ever staging of VAVFL
    - the staging is based upon an objective model derived from signal-to-noise processing, the technique of which, in itself, is novel

the modelling could also be applied to other types of visual field loss arising from drug toxicity

  - the field loss is recorded with a widely available, and regulatory authority approved, perimetric protocol
- Presents the first quantitative description of the inter-ocular mirror symmetry of VAVFL
  - the technique for quantifying symmetry has not been described before and could also be applied to other types of visual field loss
- Is based upon long-term usage of vigabatrin (median 8.8 years; IQR 5.7 to 11.3; range 0.33 to 16.1); many of these long-term exposures are unprecedented in the literature
- Contains the first illustration of a case of progressive VAVFL obtained (over a seven year period) by standard automated perimetry.
- The modelled fields from those exposed to vigabatrin are illustrated as videos in Online Resource Videos 1 2, and 3 respectively, and from the control individuals in Online Resource Video 4.

- these videos are the product of sophisticated computational methodologies
- the use of such a presentation to illustrate the appearance of the visual field is, to our knowledge, a 'first' for any journal
- the use of video material is also a first for *CNS Drugs*

**The Online Resource Video files form the basis of the manuscript and it is essential that each referee is able to view the content of these files.**

**Given the issue with one of the referees of our last manuscript, we would respectfully insist that the manuscript is NOT refereed by anyone who has previously declared a conflict of interest with Lundbeck LLC and/ or with Ovation Pharmaceuticals, Inc.**

Best wishes.

Yours Sincerely,



John Wild

**CNSA-D-19-00041**

**Objective derivation of the morphology and staging of visual field loss associated with long-term vigabatrin therapy**

We acknowledge the contribution from each of the reviewers. The review process has resulted in a much better manuscript.

All changes included in the revised manuscript are highlighted in green and are described below.

**Response to Reviewer #1**

**1. The subjects are adults. The authors have avoided discussing children but do they have any sense of whether this data would be of alarm to younger age groups.**

*We did not 'avoid(ed) discussing children'. The case series was compiled from adults on the basis of their ability to perform perimetry.*

*We have inserted a paragraph in the Discussion at the bottom of page 14/ top of page 15 of the revised manuscript.*

The model was developed from the visual fields of adults. The reduction in amplitude of the 30Hz flicker cone electroretinogram (ERG) in infants treated with vigabatrin for infantile spasms [31] is compatible with that for adults manifesting VAVFL [32]. Similarly, the topographical characteristics of the reduced peripapillary retinal nerve fibre layer thickness in children [33] is also compatible with that found in adults [34-38]. Those with vigabatrin-associated 30Hz flicker cone ERG abnormality in infancy subsequently manifest VAVFL, and corresponding retinal nerve fibre layer thinning, in later childhood/ early adolescence [39]. There is no reason to suggest, therefore, that the vigabatrin toxicity manifested in infancy will result in a different appearance to the VAVFL when the latter measure is obtained in later life.

**2. The reader would like to know, just generally, how impaired these subjects were? Were some in wheelchairs? Head control or eye movement control problem? Any with frequent seizures e.g., more than a few per day.**

*The Reviewer seems to have overlooked our commentary at the top of page 17 of the initial version of the manuscript, namely:*

All 14 individuals exhibited impaired mobility and reported symptoms explainable by their field loss, the most common of which was bumping into individuals in crowded locations.

*Nevertheless, we have modified the paragraph (now top of page 19 of the revised manuscript). The new text is given in non-italicised text.*

All 14 individuals exhibited profoundly impaired mobility and reported symptoms explainable by their VAVFL, the most common of which was bumping into individuals in crowded locations. However, some individuals with Stage 5 VAVFL also reported a negative impact on their activities of daily living.

*We have also inserted a sentence in the Methodology at the end of the paragraph on Page 7*

All individuals had been seizure free for a minimum of 24 hours prior to the visual field examination.

**3. I don't see any data on false-positive and negatives to the field tests – were there any? This makes it easier to judge the validity of the tests.**

*We made reference to the criteria for incorrect responses to the false-positive and false-negative catch trials the field tests in the initial version of the manuscript (page 7, middle paragraph).*

The reliability criteria comprised  $\leq 20\%$  incorrect responses to the fixation loss catch trials and  $\leq 15\%$  incorrect responses to the false-positive catch trials. A criterion of  $\leq 30\%$  incorrect responses was used for the false-negative catch trials although this was widened with increasing severity of the field loss [25].

*Accordingly, we have not revised the manuscript in this regard.*

**4. Do the authors have any data on visible changes to retina morphology that accompanied the field loss. And if so what stage did these develop.**

*We have inserted the following in the Discussion from the middle of page 15 onwards*

The most common bilateral finding by fundoscopy, through a dilated pupil, was either generalised or localised arteriolar narrowing which was noted at all stages of VAVFL. However, this finding was occasionally found in the presence of a normal field and, therefore, may well be associated with vigabatrin usage. Subtle bilateral retinal nerve fibre layer changes and optic nerve head pallor were noted from Stage 4 onwards. A variety of bilateral peripheral degenerative changes were also noted including hypopigmented/ white spots and surface wrinkling. However, there was considerable variation in the fundal appearance between-individuals within a given stage. The various features are in agreement with those reported previously [3, 22, 40-41] were often subtle and suggest that there is a wide spectrum of potential retinopathy associated with vigabatrin toxicity. There was no evidence of these findings in the individuals with no exposure to vigabatrin. Twenty-nine individuals exposed to vigabatrin had undergone optical coherence tomography of the peripapillary retinal nerve fibre layer at the time of the examination. All three individuals who manifested a normal visual field exhibited a normal nerve fibre layer thickness. Of the three individuals who had Stage 1 loss, two exhibited a normal thickness. An abnormally thin nerve fibre layer, characteristic of vigabatrin toxicity [34-38], was present from Stage 2 onwards in 22 individuals. The one remaining individual manifested early Stage 2 VAVFL and a normal a normal nerve fibre layer thickness.

**5. Were there any other reasons for exclusion...**

*The exclusion criteria are listed in the methods; no other exclusion criteria were applied post selection.*

**6. Do the authors have any sense of what happens if vigabatrin is stopped. Do field changes continue to progress**

*Please see the response to Query #1 from Reviewer #2. The following text has been inserted into the manuscript.*

An unpublished audit of the long-term follow-up, over a maximum of eight years, of individuals in the current study indicates that the field loss neither improves nor deteriorates following withdrawal of vigabatrin.

### Minor change

We have incorporated 'No' into the abstract.

## **Response to Reviewer #2**

### Major Comments

- 1. The major weakness of this study is failure to provide data in patients followed up longitudinally over time. Although the authors imply that patients progress from Stage 1 (least severe) to Stage 6 (most severe), they have not confirmed that such progression occurs. Most importantly, how can staging data be interpreted without knowledge of the probability of progression at each stage, and an estimate of the rate of progression? These concerns are reinforced by the finding that the occurrence of visual field defects was related to cumulative dose and duration of treatment, but there was no correlation between stages and exposure. The authors' comment that toxicity is 'idiosyncratic' does not explain the lack of correlation between severity of the defect and dose/duration of exposure. These issues need to be discussed.**

*We have inserted an essentially new paragraph in the Discussion on Page 17. The new text is given in non-italicised text.*

*The staging of VAVFL was derived from cross-sectional evidence and does not imply progressive loss. Due to the potency of the potential toxicity, most individuals had been withdrawn from vigabatrin either immediately prior to the introduction of the visual field examination into the care regime or following confirmation of the VAVFL. As such, neither the probability of progression at each stage nor an estimate of the rate of progression can be determined. However, the presumption is that, given continued vigabatrin therapy, VAVFL will progress through the various stages. The lack of an association between the stage of loss, at detection, and either the duration or the cumulative dose of vigabatrin implies that the relationship between the extent of exposure and both the onset and the severity of VAVFL varies depending upon the individual susceptibility to vigabatrin. However, the time of detection is not the time to onset of the VAVFL. The rate of any subsequent progression, therefore, remains unknown. A case of progressive loss during approximately 7.75 years of vigabatrin therapy and illustrated in terms of the Pattern Deviation probability map of the Central 24-2 Threshold Test and, subsequently, the C30-2T is shown in Fig 6. The outcome of the corresponding FF135 at the final visit is shown in Online Resource Fig 4. VAVFL has also been shown to be progressive whilst on therapy in all 14 cases over a mean follow-up of 10.7 years with the rate of reduction in the 14e isopter of kinetic perimetry increasing with increase in cumulative dose [6]. An unpublished audit of the long-term follow-up, over a maximum of eight years, of individuals in the current study indicates that VAVFL neither improves nor deteriorates following withdrawal of vigabatrin.*

## **2. Selection bias**

*We have inserted a sentence relating to selection bias in the last sentence on Page 6.*



There was no conscious selection bias. Patients were included on the basis of sequential retrieval of cases which met the inclusion/ exclusion criteria.

### **3. Continuity in severity of field loss across stages. How were cutoffs across stages (and the actual number of stage) defined?**

*We have modified the paragraph (now at the bottom of page 9/ top of page 10 of the revised manuscript). The new text is given in non-italicised text.*

*The stages were empirically selected on the basis of clinically meaningful intervals of peripheral field loss and, subsequently, of central field loss. These intervals, in turn, were based upon the magnitude of the between-examination physiological variability in the differential light sensitivity as a function of severity of loss [28-29]. The derivation ensured continuity across stages.*

### **4. Please clarify whether the ophthalmologist was aware of the exposure status.**

*We have inserted a commentary on page 15.*

The ophthalmological examinations were undertaken by any one of four ophthalmologists, depending upon the particular clinic, who were all unaware of the findings from their colleagues. All four were aware of the anti-epileptic drug history and, usually, of the visual field. However, the modelling of the visual fields from those exposed to vigabatrin and from the control individuals was objective and independent of the outcome from the ophthalmological and neurological examinations.

### **Compile Table about characteristics/ Statistical differences**

*We have inserted the summary statistics for the duration and cumulative dose of vigabatrin into the penultimate paragraph on page 12 of the revised manuscript.*

Those with VAVFL manifested a greater cumulative dose (mean 7.94kg; SD 4.45; range 1.1 to 20.7) and a longer duration of therapy (8.86 years; SD 3.51; range 0.66 to 16.05) at the time of perimetry than those without field loss (mean 3.36kg; SD 4.77; range 0.11 to 16.14; and mean 3.62 years; SD 3.94; range 0.33 to 11.83; respectively). The difference between means (Student's t-test for two independent samples) were 4.58 kg (95% CI 1.71-7.45;  $p < 0.001$ ) and 5.25 years (95% CI 2.98-7.53;  $p < 0.001$ ), respectively.

*We have inserted and additional sentence on Page 12.*

There was no association between the stage of VAVFL and either age, age at onset of epilepsy or age at onset of vigabatrin.

### **Minor Comments**

#### **1. Remove without prescription**

*The statement concerning the free availability on-line without prescription has been removed from the revised manuscript*

**2. Lines 26-50 This paragraph does not describe appropriately the background for the study.**

*The Reviewer is incorrect in his/ her commentary.*

- i) *The best description of the methodology used to compile the Registry described by Krauss et al (2016) is that of Pellock et al (2011) [Epilepsy & Behavior 2011;22:710-717].*

*Page 2 Paragraph 4 of Pellock et al (2011) states that:*

*'Regular ophthalmologic assessments are required throughout vigabatrin therapy: at baseline ( $\leq 4$  weeks after therapy initiation), at least every 3 months during therapy and 3 to 6 months after discontinuation. Fulfilment of this mandate is documented by registry receipt of an ophthalmologic assessment form.....In many instances, the first actual vision test may be conducted well after entry into the registry.'*

*The Registry was an FDA mandated requirement for the marketing authorisation of Sabril within the USA. The data analysis for the Krauss et al (2016) paper was undertaken by a third party institution and the editorial support for the paper was funded by the Marketing Authorisation Holder for Sabril in the USA. The paper is carefully worded, is ambiguous in crucial places, and largely concentrates on the number of individuals enrolled within the Registry and the likelihood of retention of these individuals over time. The referee is correct in his/ her assertion that the paper by Krauss et al (2016) 'does not provide (sic) baseline data on visual fields'; however, the paper states that:*

*'Clinicians reported that eight patients discontinued treatment because of visual field defects (VFD). The patients ranged from ages 18 to 55 years at entry into the SHARE program and were divided equally by gender. One 19 year old patient was reported to have an indication of IS, the other patients had CPS. Three patients entered the program with previous vigabatrin exposure. The five patients who were newly treated had vigabatrin discontinued by their clinicians because of VFD after exposures of 13 months to 3.3 years (Mean: 23 months).'*

*The clear implication from the above is that the visual field loss in at least five patients occurred after the baseline examination.*

*There is no discussion in the paper on the (low) prevalence of visual field loss and no mention of the fact that 'the data are not interpreted as indicating lack of visual toxicity.'*

*Although not stated, the clear implication to the reader is that the prevalence of the toxicity, derived from a [this] prospective study, is much lower than that derived from the (European) retrospective studies.*

- ii) *The Sergott et al (2016) paper is described as a 'small study sponsored by Lundbeck.' It should be noted that this study was a one year prospective study was mandated by the FDA.*

*Given the above and given the acceptance by Reviewer #1, we have not altered the paragraph.*

## **Response to Comments from Editor**

- 1. We have moved the definition of VAVFL to the end of the figure legend*
- 2. We have renumbered the figures as advised*
- 3. We have revised the format of the supplementary content as advised*

*Our preference would be to embed a link to the videos within the article.*

## Staging of vigabatrin-associated visual field loss

**Objective derivation of the morphology and staging of visual field loss associated with long-term vigabatrin therapy**

John M Wild<sup>1</sup>, Phillip E M Smith<sup>2</sup>, Carlo Knupp<sup>1</sup>

<sup>1</sup> College of Biomedical Sciences, Cardiff University, Maindy Road, Cardiff CF24 4HQ, United Kingdom

<sup>2</sup> Alan Richens Unit, Welsh Epilepsy Centre, University Hospital of Wales, Heath Park, Cardiff, CF14 4XW, United Kingdom.

**Corresponding Author:** John Wild.

**Address:** College of Biomedical Sciences, Cardiff University, Maindy Road, Cardiff CF24 4HQ, United Kingdom.

**Telephone:** ++44 29 20 87 64 87

**Fax:** ++44 29 20 87 48 59

**Email:** [wildjm@cardiff.ac.uk](mailto:wildjm@cardiff.ac.uk)

**John Wild:** ORCID: 0000-0003-3019-3889

**Running head:** Staging of vigabatrin-associated visual field loss

**Enhanced digital features**

To view enhanced digital features for this article go to <https://doi.org/10.6084/m9.figshare.7981592>

## Abstract

**Background** The morphology and between-eye symmetry of the visual field loss associated with the anti-epileptic drug vigabatrin (VAVFL) has received little attention.

**Objective** To model the appearance, and ensuing staging, of VAVFL, derived with the European Medicines Agency approved perimetric protocol.

**Methods** A retrospective, cross-sectional, observational design identified 123 adults who had received vigabatrin for refractory seizures and who had no evidence of co-existing retinogeniculo-cortical visual pathway abnormality. Thirty-eight adults with refractory seizures and identical inclusion criteria, but no exposure to vigabatrin, acted as controls. For each group, the median outcome at each stimulus location in each eye (of absolute loss, relative loss or Pattern Deviation probability level, as appropriate) was derived for each successive ten pairs of fields, ranked for severity. Between-eye symmetry was quantified by an index which accounted for severity of loss and which was referenced to the likelihood of the occurrence of symmetry due to chance.

**Results** The modelled VAVFL was bilateral and highly symmetrical and was described by six stages which were all independent of the extent of vigabatrin exposure. The loss originated in the extreme temporal periphery and encroached centripetally along all meridians towards fixation. The initial appearance within the central field (Stage Two) occurred inferior-nasally. Subsequent stages exhibited increasing loss which was greater nasally than temporally. Stage Six described concentric loss extending to approximately 15° eccentricity from fixation.

**Conclusion** The model exhibited a consistent pattern of VAVFL. The staging of the loss could assist the risk:benefit analysis of vigabatrin for the treatment of epilepsy.



## Key Points

In this study of 123 individuals treated with vigabatrin as adjunct therapy for refractory focal seizures and without evidence of other retino-geniculo-cortical visual pathway abnormality, objective modelling, based upon convolution theory used in signal-to-noise processing, identified a consistent pattern of bilateral symmetrical visual field loss obtained using a regulatory approved perimetric protocol. No modelled field loss was present for the 38 individuals with identical inclusion criteria but no exposure to vigabatrin.

Six stages of modelled vigabatrin-associated field loss (VAVFL) were evident. Originating in the extreme temporal periphery, the field loss encroached, with increasing severity, centripetally along all meridians towards fixation with greater nasal than temporal loss. Stage Six manifested as concentric loss to within approximately 15° from fixation.

The between-eye mirror symmetry of the VAVFL at all stages, quantified by a novel index, which accounted for the severity of loss, was not attributable to chance ( $p \leq 0.01$ ).

The use of either the Central 30-2 or the Central 24-2 Threshold Tests, alone, will not identify VAVFL until late Stage Two and late Stage Three, respectively.

## 1.0 Introduction

The anti-epileptic drug vigabatrin has long been associated with visual field loss (VAVFL) [1-2] which manifests as a bilateral concentric peripheral defect over a continuum of severity and which, generally, also involves the central field to varying extents [3-4]. The field loss is considered to be slowly progressive [5-7] irreversible but non-progressive on withdrawal from vigabatrin [8-9] and asymptomatic until severe loss is present [3, 10-12]. The modelled prevalence of VAVFL in adults increases rapidly after the first two years (2kg cumulative dose) of exposure [13-14] and can reach 75-80% after approximately six years (5kg cumulative dose) [14]. Vigabatrin is now available generically.

The association between vigabatrin and ocular toxicity is currently being re-evaluated [15-17] on the basis that the attribution of the toxicity emanates from retrospective studies which do not include a pre-treatment baseline evaluation. The lack of evidence for the toxicity in the two prospective studies alongside the finding that approximately 25% of individuals suitable for vigabatrin exhibited clinically significant pre-existing reduced visual acuity and/ or visual field loss and/ or peripapillary retinal nerve fibre layer thinning, has led to the suggestion that abnormalities of the afferent system are a co-morbidity of severe refractory epilepsy and are unrelated to vigabatrin [15-16]. However, the limited exposures to vigabatrin and the poor retention of individuals, within these two studies [15-16], together with inappropriate visual field methodologies and analyses, have not facilitated detection of the toxicity.

Surprisingly, there has been little emphasis on the areal and depth characteristics, the between eye-symmetry, and the staging, of VAVFL. Such information is essential not only to identify the

onset and the progression of the VAVFL but also to differentiate it from other types of (concomitant) field loss.

The areal extent and the depth of any type of visual field loss is highly dependent upon the perimetric technique used for the examination. VAVFL has been investigated by a variety of methods [7, 10-11, 16, 18-23] but the characteristics of the field loss by each technique have received little attention. Unfortunately, due to the peripheral nature of the field loss, there is no one ideal technique, across the current range of commercially available perimeters, for the assessment of VAVFL. The most pragmatic approach is that approved by the European Medicines Agency (EMA) [23] namely, Three Zone (two luminance level) suprathreshold perimetry, referenced to the age-corrected threshold, of the peripheral field out to a maximum temporal extent of 90° eccentricity and standard automated perimetry of the central field (SAP).

Given the on-going re-evaluation of the association between vigabatrin and visual field loss and the wider availability of the drug, a description of the morphology and staging of the VAVFL resulting from the EMA approved protocol, and derived from individuals with long-term exposures to vigabatrin, would be of benefit to clinicians and to patients, alike. If such a description was based upon a technique which reduced the signal-to-noise ratio and was derived from individuals with no evidence of retino-geniculo-cortical visual pathway abnormality, except that of potential vigabatrin toxicity, the characteristics of the field loss attributable to vigabatrin would become manifest from backgrounds of any unexpected field loss and of any increased variability associated with the response by the patient. Such an approach would also enable the use of the visual fields from individuals with long-term exposures to vigabatrin but with no pre-treatment baseline.

The aim of the study, therefore, was threefold: to model the morphology of VAVFL obtained with the EMA approved perimetric protocol and resulting from long-term vigabatrin therapy; to quantify any between-eye symmetry; and to describe the ensuing staging of the field loss.

## 2.0 Methods

### 2.1 Case series

A retrospective case series comprising 161 adults with refractory complex partial (focal) seizures was compiled from those presenting to the Alan Richens Unit of the Welsh Epilepsy Centre, University Hospital of Wales, Cardiff, UK, and to associated clinics. All individuals had undergone part or all of the EMA approved protocol with the Humphrey Field Analyzer 750 (Carl Zeiss, Meditec, Dublin, CA) namely, the Full Field 135 Point Test with the three zone age-corrected strategy (FF135) and the Central 30-2 Threshold Test (C30-2T) with the FASTPAC strategy. None exhibited ocular or intra-cranial conditions resulting in, or likely to cause, visual field loss, as determined by ophthalmological and neurological examination including whole-brain magnetic resonance imaging. Further exclusion criteria comprised the presence of structural and/or functional ocular abnormalities likely to impede or confound the outcome of perimetry and which are described elsewhere [24]. There was no conscious selection bias. Individuals were included on the basis of sequential retrieval of cases which met the inclusion/exclusion criteria.

## 2.2 Vigabatrin exposure

Of the 161 individuals (Fig 1, top row), 123 had been exposed to vigabatrin as add-on therapy. The remaining 38 individuals had received a variety of other antiepileptic drugs and were included as age-matched controls.

## 2.3 Perimetry

Sixty-three of the 123 individuals exposed to vigabatrin and all 38 of those exposed to other antiepileptic drugs had undergone the complete EMA approved protocol (Fig 1, second row). Of the remaining 60 individuals exposed to vigabatrin, 24 had undergone the FF135, only, and 36 had undergone the C30-2T, only (Fig 1, second row) as ordered by the treating neurologist on clinical grounds. Thus, 87 of the 123 individuals exposed to vigabatrin had undertaken the FF135 and 99 individuals had undertaken the C30-2T.

The visual fields from the first occasion, after the first visit, at which reliable perimetric outcomes had been achieved were used for the individuals in each of the two groups. The reliability criteria comprised  $\leq 20\%$  incorrect responses to the fixation loss catch trials and  $\leq 15\%$  incorrect responses to the false-positive catch trials. A criterion of  $\leq 30\%$  incorrect responses was used for the false-negative catch trials although this was widened with increasing severity of the field loss [25]. All individuals had been seizure free for a minimum of 24 hours prior to the visual field examination.



## 2.4 Morphology of visual field loss

The Single Field Printouts of the FF135 and of the C30-2T were exported to a personal computer via a Humphrey Field Analyzer 820 and saved in Tagged Image File Format. The outputs from each eye for both types of perimetry were then read by custom Naïve Bayesian character recognition software [26].

The fields were separately sorted for each group, and for each type of perimetry, as right and left eye pairs, and ranked from the least to the most affected field. The severity of the given pair of fields was defined as the sum, across the two eyes, of the number of defects, weighted to account for relative or absolute loss, respectively, derived by the FF135 and/ or for the probability level of the Pattern Deviation probability map derived by the C30-2T. In the case of those undergoing both tests, the outcome from the C30-2T was more heavily weighted.

The median outcome for each group at each stimulus location in each eye for each type of perimetry was then separately calculated for each successive ten pairs of fields, ranked for severity. Such an approach was based upon convolution theory used in signal processing [27].

The procedure resulted in 54 pairs of modelled fields from the 63 individuals exposed to vigabatrin who had undergone the combined EMA protocol; 78 pairs for the FF135 and 90 for the C30-2T. Twenty-nine pairs of modelled fields resulted from the 38 individuals with no vigabatrin exposure. The outcomes for the right and left eyes, separately, were then displayed in terms of Audio Video Interleaved (AVI) movies.

## 2.5 Between-eye mirror symmetry of visual field loss

The between-eye mirror symmetry of the field was expressed in terms of a single index which comprised two components, each of which was a fraction. The index accounted for the severity of loss and was compared to the likelihood of obtaining symmetrical field loss due to chance.

The numerator and denominator of the first component constituted of, respectively, the number of mirror image stimulus locations between the two eyes which exhibited either relative or absolute loss for the FF135, or abnormality at  $p \leq 0.02$  by Pattern Deviation probability analysis for the C30-2T, and the total number of locations exhibiting abnormality across the two eyes.

The numerator and denominator of the second component comprised, respectively, the total number of locations exhibiting abnormality across the two eyes and the total number of stimulus locations across the two eyes. The index was compared to that obtained from the simulation of one million randomly generated pairs of abnormal fields which manifested varying locations, areas and depths of field loss both within- and between-eyes. The concept was such that symmetry would become apparent when compared to a series of pairs of fields with varying characteristics of asymmetric field loss.

The symmetry index was separately calculated for each pair of peripheral, and each pair of central fields, for the modelled fields and for the measured fields.

## 2.6 Staging of the measured fields for those exposed to vigabatrin

The staging of the VAVFL was derived from the modelled fields. The stages were empirically selected on the basis of clinically meaningful intervals of peripheral field loss and, subsequently, of central field loss. These intervals, in turn, were based upon the magnitude of the between-examination physiological variability in the differential light sensitivity as a function of severity

of loss [28-29]. The derivation ensured continuity across stages. The staging of the modelled peripheral fields was undertaken masked to that of the modelled central fields. The outcome of the staging of the central and peripheral fields, with respect to one another, was then validated based upon the outcome of the model from the individuals who had undergone the combined protocol. The staging was then applied to each of the 87 pairs of measured fields from the FF135 and to each of the 99 measured fields from the C30-2T.

### 3.0 Results

#### 3.1 Demographic characteristics of the case series

Of the 123 individuals exposed to vigabatrin, 50 were male and 73 female. Thirty-five were receiving vigabatrin at the time of perimetry. Of the 38 age-matched controls, 17 were male and 21 female. The mean ages of the two groups, at the time of perimetry, were 40.8 years (SD 13.6) and 38.5 years (SD 10.0), respectively.

#### 3.2 Morphology of VAVFL

The rolling medians of the 78 pairs of modelled fields for the FF135 from those exposed to vigabatrin; of the 90 pairs for the C30-2T; and of the 54 pairs for the combined EMA protocol are given in video format in Online Resource Figs 1, 2 and 3, respectively (available from URL: <https://doi.org/10.6084/m9.figshare.7981592>).

The rolling medians of the 29 pairs of modelled fields for the combined EMA protocol from those with no exposure to vigabatrin are given in video format in Online Resource Fig 4

(available from URL: <https://doi.org/10.6084/m9.figshare.7981592>). The modelled fields showed was no evidence of visual field loss.

### 3.3 Staging of VAVFL

The staging of the FF135 and of the C30-2T modelled fields for those exposed to vigabatrin is shown in Figs 2 to 4.

Sixteen of the 36 individuals exposed to vigabatrin, who had undergone the C30-2T, only, manifested a normal measured central field (Fig 1 third row). These 16 fields were included in the model, but, in the absence of a peripheral field assessment, the individuals could not be categorised as to the potential outcome of the toxicity and were considered as equivocal.

Of the remaining 107 individuals exposed to vigabatrin (Fig 1, fourth row), 12 exhibited normal measured fields and 95 field loss which conformed to the modelled fields and which was therefore designated as VAVFL.

The frequency, by stage, of VAVFL is shown in Fig 1 (bottom row). Fourteen of the 95 individuals with VAVFL exhibited Stage One VAVFL, 21 Stage Two, 14 Stage Three, 22 Stage Four, 10 Stage Five and 14 Stage Six.

The relationship between the stage of loss and the summary measures of the visual field, Mean Deviation (MD) and Pattern Standard Deviation, for the C30-2T measured fields is shown in Fig 5.

### 3.4 Between-Eye Mirror Symmetry of VAVFL

The two components of the symmetry index for the FF135 and for the C30-2T are shown in Online Resource Videos 1, 2 and 3; at each stage of the modelled VAVFL in Figs 2 to 4; and for each measured field of those exposed to vigabatrin in Fig 6.

### 3.5 Demographics of VAVFL

The difference in the proportion with VAVFL by gender, 39 out of 44 males and 56 out of 63 females, was not statistically significant.

The mean age, at the time of perimetry, of those with VAVFL, 41.6 years (SD 14.4), was also similar to that of those exposed to vigabatrin but with normal fields, mean 37.9 years (SD 7.9).

Those with VAVFL manifested a greater cumulative dose (mean 7.94kg; SD 4.45; range 1.1 to 20.7) and a longer duration of therapy (8.86 years; SD 3.51; range 0.66 to 16.05) at the time of perimetry than those without field loss (mean 3.36kg; SD 4.77; range 0.11 to 16.14; and mean 3.62 years; SD 3.94; range 0.33 to 11.83; respectively). The difference between means (Student's t-test for two independent samples) were 4.58 kg (95% CI 1.71-7.45;  $p<0.001$ ) and 5.25 years (95% CI 2.98-7.53;  $p<0.001$ ), respectively.

There was no evidence of a relationship between the stage of the VAVFL and the extent of the exposure to vigabatrin at the time of perimetry. The median exposures at the time of perimetry which resulted in Stage One loss were 7.7kg cumulative dose (IQR 4.8, 12.1; range 1.5-13.3) and 9.4 years (IQR 6.2, 12.1; range 2.4-16.1) and which resulted in Stage 6 loss were 6.7kg cumulative dose (IQR 4.4, 10.6; range 2.8-19.0) and 7.7 years (IQR 5.3, 10.6; range 3.1-13.0).

There was also no association between the stage of VAVFL and either age, age at onset of epilepsy or age at onset of vigabatrin.

### 4.0 Discussion

The objective rolling median and symmetry index outcomes from the FF135 and the C30-2T confirm that vigabatrin is associated with symmetrical bilateral visual field loss. The VAVFL manifests over a range of severities that can be described in six stages. The field loss originates in the extreme temporal periphery and, as the severity increases, encroaches centripetally along all meridians towards fixation resulting in bi-nasal loss and relative temporal sparing until the end stage (Stage Six) which manifests as a concentric loss to within approximately 15° from fixation. The initial appearance in the central field (Stages Two and Three) is primarily inferior-nasally. VAVFL exhibits a high degree of between-eye mirror symmetry at each stage of loss. In contrast, the modelled field for the age-matched control individuals with no exposure to vigabatrin was entirely normal.

The apparent absence of initial loss superior-nasally within the central field is most likely attributable to the greater defect depth in this region required to achieve statistical significance. The latter, in turn, arises from the wider distribution of normal values arising from the between-individual variation in the position of the upper eyelid and from the increased variability in response associated with the steeper gradient of the superior visual field, compared to those inferior-nasally.

The fundamental strengths of this retrospective study were the robust exclusion of individuals with retino-geniculo-cortical visual pathway abnormality; the inclusion of long-term exposures to vigabatrin (median 8.8 years; IQR 5.7 to 11.3; range 0.1 to 16.1) many of which are unparalleled in the literature; and the utilisation of the two novel objective techniques which have not previously been applied to perimetry. These latter techniques enabled objective descriptions of the characteristics and between-eye mirror symmetry of VAVFL.

The modelled field at all six stages is in agreement with that derived from cross-sectional retrospective evidence with kinetic perimetry [4]. The initial manifestation of the VAVFL in this latter model was defined as a ‘non-seen’ response to the Goldmann V4e isopter at 80° temporally and/ or at 40° nasally. This definition is entirely consistent with the location and depth of the loss described as Stage One in the current study: the absolute loss designated by the FF135 is equivalent to the Goldmann V4e stimulus. The increasing encroachment into the nasal field with relative sparing of the temporal field, during Stages Two to Five, is compatible with the model out to approximately 80° temporally, derived from cross-sectional evidence, using suprathreshold perimetry equivalent to the I4e isopter [30].

The model was developed from the visual fields of adults. The reduction in amplitude of the 30Hz flicker cone electroretinogram (ERG) in infants treated with vigabatrin for infantile spasms [31] is compatible with that for adults manifesting VAVFL [32]. Similarly, the topographical characteristics of the reduced peripapillary retinal nerve fibre layer thickness in children [33] is also compatible with that found in adults [34-38]. Those with vigabatrin-associated 30Hz flicker cone ERG abnormality in infancy subsequently manifest VAVFL, and corresponding retinal nerve fibre layer thinning, in later childhood/ early adolescence [39]. There is no reason to suggest, therefore, that the vigabatrin toxicity manifested in infancy will result in a different appearance to the VAVFL when the latter measure is obtained in later life.

The ophthalmological examinations were undertaken by any one of four ophthalmologists, depending upon the particular clinic, who were all unaware of the findings from their colleagues. All four were aware of the anti-epileptic drug history and, usually, of the visual field outcome. However, the modelling of the visual fields from those exposed to vigabatrin and from the control individuals was objective and independent of the outcome from the ophthalmological and neurological examinations.

The most common bilateral finding by fundoscopy, through a dilated pupil, was either generalised or localised arteriolar narrowing which was noted at all stages of VAVFL. However, this finding was occasionally found in the presence of a normal field and, therefore, may well be associated with vigabatrin usage. Subtle bilateral retinal nerve fibre layer changes and optic nerve head pallor were noted from Stage 4 onwards. A variety of bilateral peripheral degenerative changes were also noted including hypo-pigmented/ white



spots and surface wrinkling. However, there was considerable variation in the fundal appearance between-individuals within a given stage. The various features are in agreement with those reported previously [3, 22, 40-41], were often subtle, and suggest that there is a wide spectrum of potential retinopathy associated with vigabatrin toxicity. There was no evidence of such findings in the individuals with no exposure to vigabatrin. Twenty-nine individuals exposed to vigabatrin had undergone optical coherence tomography of the peripapillary retinal nerve fibre layer at the time of the examination. All three individuals who manifested a normal visual field exhibited a normal nerve fibre layer thickness. Of the three individuals who had Stage 1 loss, two exhibited a normal thickness. An abnormally thin nerve fibre layer, characteristic of vigabatrin toxicity [34-38], was present from Stage 2 onwards in 22 individuals. The one remaining individual manifested early Stage 2 VAVFL and a normal a normal nerve fibre layer thickness.

The outcome of the current study and of the earlier models [4, 28] indicates that perimetry out to 60° eccentricity [42] is inadequate to detect Stage One and also possibly early Stage Two loss. Similarly, a worsening of 3dB from the baseline Mean Deviation (MD) index for the C30-2T, which was used as a primary outcome measure for the presence of VAVFL in the prospective Phase IV study [16], will not detect VAVFL until late in Stage Two or early in Stage Three. The MD is a summary measure of the central field and does not describe the spatial appearance of visual field loss: in the current study, the median MDs for the individuals exposed to vigabatrin and for the controls was -1.75dB and -1.23dB, respectively.

It is clear from the staging of the VAVFL that perimetry should initially be undertaken with, in the case of the Humphrey Field Analyzer, the FF135. If the field is normal to the FF135 or

exhibits Stage One abnormality, the C30-2T is unnecessary. If the peripheral field exhibits Stage Two abnormality to the FF135, the C30-2T should also be undertaken to reveal the full depth of the loss since the magnitude of the Pattern Deviation value required for statistical significance at the extreme nasal locations within the central field can be less than the 8dB suprathreshold increment of the stimulus luminance of the FF135. If the FF135 indicates both peripheral and central loss, i.e., Stage Three and worse, subsequent examinations need only be undertaken with the C30-2T. The Central 24-2 Threshold Test, alone, should only be used for Stage Six. The characteristics of VAVFL are more prominent with the FASTPAC algorithm than with the SITA Standard or Fast algorithms which were designed to detect glaucomatous field loss. The peripheral manifestation of VAVFL is fortuitous since, in our experience, patients exposed to vigabatrin find suprathreshold perimetry considerably easier to perform than SAP. In cases where patients are unable to undertake SAP, assessment with the FF135, although collecting less information about relative loss, is acceptable.

The staging of VAVFL was derived from cross-sectional evidence and does not imply progressive loss. Due to the potency of the potential toxicity, most individuals had been withdrawn from vigabatrin either immediately prior to the introduction into the care regime of the visual field examination or following confirmation of the VAVFL. As such, neither the probability of progression at each stage nor an estimate of the rate of progression can be determined. However, the presumption is that, given continued vigabatrin therapy, VAVFL will progress through the various stages. The lack of an association between the stage of loss, at detection, and either the duration or the cumulative dose of vigabatrin implies that the relationship between the extent of exposure and both the onset and the severity of VAVFL varies depending upon the individual susceptibility to vigabatrin. However, the time of detection is not the time to onset of the VAVFL. The rate of any subsequent progression,

1 therefore, remains unknown. A case of progressive loss during approximately 7.75 years of  
2 vigabatrin therapy and illustrated in terms of the Pattern Deviation probability map of the  
3 Central 24-2 Threshold Test and, subsequently, the C30-2T is shown in Fig 6. The outcome  
4 of the corresponding FF135 at the final visit is shown in Online Resource Fig 4. Progressive  
5 loss whilst on therapy has been shown over a mean follow-up of 10.7 years; the rate of  
6  
7 reduction in the I4e isopter of kinetic perimetry increased with increase in cumulative dose  
8  
9 [6]. An unpublished audit of the long-term follow-up, over a maximum of eight years, of  
10  
11 individuals in the current study indicates that the VAVFL neither improves nor deteriorates  
12  
13 following withdrawal of vigabatrin.  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24

25 The appearance of Stage One VAVFL should act as an alert signal for a re-evaluation of the  
26 management of the epilepsy. This may encompass either a change in anti-epileptic drug  
27 therapy or an increase in the frequency of neuro-ophthalmological monitoring. However, the  
28 extreme peripheral stimulus locations of the FF135 are also influenced by the facial anatomy,  
29 particularly superiorly and inferior-nasally. The outcome of the examination at such  
30 locations, and those in the extreme temporal periphery, is also dependent upon the vigilance  
31 of the patient. Indeed, normal individuals do not necessarily produce statistically normal  
32 visual fields. The FF135 commences with the examination of the central field. Once  
33 completed, the examination pauses for removal of the trial lenses prior to the examination of  
34 the peripheral field. Instructions to the patient, during this pause, to the effect that the stimuli  
35 are about to appear in the (extreme) periphery, together with an exhortation to keep the eye  
36  
37 ‘wide open’, generally leads to a ‘seen’ response, in the normal eye, at these extreme  
38 locations. However, individuals with epilepsy frequently exhibit greater within-examination  
39 variability than normal individuals, which can lead to the impression of visual field loss, and  
40 approximately 30% of individuals with epilepsy are unable to undertake perimetry of any  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60  
61  
62  
63  
64  
65

type [15, 32]. The clinical skill lies in the identification of actual visual field loss, characteristic of the given lesion, from apparent visual field loss due to increased variability and/ or an inability to understand the requirements of the test. In addition, VAVFL frequently co-exists with that arising from an intra-cranial lesion.

Fourteen individuals exhibited Stage Six VAVFL. Great care was taken to exclude individuals manifesting visual field loss of suspected psychogenic origin from the case series [44]. Three of these 14 individuals had undergone optical coherence tomography and exhibited severe thinning of the retinal nerve fibre layer. All 14 individuals exhibited profoundly impaired mobility and reported symptoms explainable by their VAVFL, the most common of which was bumping into individuals in crowded locations. However, some individuals with Stage 5 VAVFL also reported a negative impact on their activities of daily living.

The study utilised a retrospective cross-sectional design. However, the objective derivation of field loss based upon signal-to-noise theory, further supported by the stringent ophthalmological and neurological inclusion criteria, uniquely facilitated a retrospective design and largely obviated the need for a prospective study referenced to a pre-treatment baseline.

The variable and lengthy time to onset of VAVFL together with the opportunity to study a wide range of long-term exposures, further highlights the pragmatism of this retrospective study compared to a prospective study. Nevertheless, prospective longitudinal studies over

the long-term would unequivocally rule out the presence of abnormality at baseline and would also determine the evolution of the field loss.

## 5.0 Conclusions

The appearance of VAVFL has received little attention. This absence of knowledge hinders not only the recognition of the onset, and of any progression, of the field loss but also the differentiation from other (concomitant) types of loss. The staging of the VAVFL derived from a regulatory approved perimetric protocol, and reported here, should assist clinicians and patients, alike, in the risk:benefit analysis of vigabatrin for the treatment of epilepsy.

## 6.0 Compliance with Ethical Standards

**Funding** None

**Conflicts of interest** John M Wild, Phillip EM Smith and Carlo Knupp declare that they have no conflicts of interest.

**Ethical approval** All procedures performed in studies involving human participants were in accordance with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. The Local Research and Ethics Committee ruled that approval was not required for this study. The visual field assessments were considered to be part of normal good clinical practice. The visual fields were de-identified and so written informed consent was not required.

**Data availability** The datasets generated and/or analysed during the current study are available from the corresponding author on reasonable request.

## 7.0 References

1. Eke T, Talbot JF, Lawden MC. Severe persistent visual field constriction associated with vigabatrin. *BMJ*. 1997;314(7075):180-1.
2. Maguire MJ, Hemming K, Wild JM, et al. Prevalence of visual field loss following exposure to vigabatrin therapy: a systematic review. *Epilepsia*. 2010;51(12):2423-31.
3. Wild JM, Martinez C, Reinshagen G, Harding GF. Characteristics of a unique visual field defect attributed to vigabatrin. *Epilepsia*. 1999;40(12):1784-94.
4. Malmgren K, Ben-Menachem E, Frisén L. Vigabatrin visual toxicity: evolution and dose dependence. *Epilepsia*. 2001;42(5):609-15.
5. Hardus P, Verduin W, Postma G, et al. Long term changes in the visual fields of patients with temporal lobe epilepsy using vigabatrin. *Br J Ophthalmol*. 2000;84(7):788-90.
6. Clayton LM, Stern WM, Newman WD, et al. Evolution of visual field loss over ten years in individuals taking vigabatrin. *Epilepsy Research*. 2013;105(3):262-71.
7. Nowomiejska K, Jedrych M, Brzozowska A, et al. Relationship between the area of isopters and vigabatrin dosage during two years of observation. *BMC Ophthalmol*. 2014;14:56. doi: 10.1186/1471-2415-14-56.
8. Johnson MA, Krauss GL, Miller NR, et al. Visual function loss from vigabatrin: effect of stopping the drug. *Neurology*. 2000;55(1):40-5.

9. Nousiainen I, Mäntyjärvi M, Kälviäinen R. No reversion in vigabatrin-associated visual field defects. *Neurology*. 2001;57(10):1916-7.
10. Daneshvar H, Racette L, Coupland SG, et al. Symptomatic and asymptomatic visual loss in patients taking vigabatrin. *Ophthalmology*. 1999;106(9):1792-8.
11. Kälviäinen R, Nousiainen I, Mäntyjärvi M, et al. Vigabatrin, a gabaergic drug, causes concentric visual field defects. *Neurology*. 1999;53(5):922-6.
12. Hardus P, Verduin WM, Postma G, et al. Concentric contraction of the visual field in patients with temporal lobe epilepsy and its association with the use of vigabatrin medication. *Epilepsia*. 2000;41(5):581-7.
13. European Medicines Agency. Opinion of the Committee for proprietary medicinal products pursuant to Article 12 of Council Directive 75/319/EEC as amended for vigabatrin. Annex 1. Scientific conclusions and grounds for amendment of the summaries of product characteristics presented by the EMEA. 1999, European Medicines Agency, Canary Wharf, London, United Kingdom. Available at:  
[https://www.ema.europa.eu/documents/referral/opinion-committee-proprietary-medicinal-products-pursuant-article-12-council-directive-75/319/eec-amended-vigabatrin-annexes-i-ii-iii-iv\\_en.pdf](https://www.ema.europa.eu/documents/referral/opinion-committee-proprietary-medicinal-products-pursuant-article-12-council-directive-75/319/eec-amended-vigabatrin-annexes-i-ii-iii-iv_en.pdf) Accessed 6th April 2019.
14. Wild JM, Fone DL, Aljarudi S, et al. Modelling the risk of visual field loss arising from long-term exposure to the anti-epileptic drug vigabatrin: a cross-sectional approach. *CNS Drugs*. 2013;27(10):841-9.
15. Krauss G, Faught E, Foroozan R, et al. Sabril® registry 5-year results: Characteristics of adult patients treated with vigabatrin. *Epilepsy Behav*. 2016;56(3):15-19.

16. Sergott RC, Johnson CA, Laxer KD, et al. Retinal structure and function in vigabatrin-treated adult patients with refractory complex partial seizures. *Epilepsia* 2016;57(10):1634-42.
17. Foroozan R. Vigabatrin: lessons learned from the United States experience. *J Neuro-Ophthalmol.* 2018;38(4):442-50.
18. Hardus P, Verduin WM, Engelsman M, et al. Visual field loss associated with vigabatrin: quantification and relation to dosage. *Epilepsia.* 2001;42(2):262-7.
19. Kinirons P, Cavalleri GL, O'Rourke D, et al. Vigabatrin retinopathy in an Irish cohort: lack of correlation with dose. *Epilepsia.* 2006;47(2):311-7.
20. Manuchehri K, Goodman S, Siviter L, Nightingale S. A controlled study of vigabatrin and visual abnormalities. *Br J Ophthalmol.* 2000;84(5):499-505.
21. van der Torren K, Graniewski-Wijnands HS, Polak BC. Visual field and electrophysiological abnormalities due to vigabatrin. *Doc Ophthalmol.* 2002;104(2):181-8.
22. Miller NR, Johnson MA, Paul SR, et al. Visual dysfunction in patients receiving vigabatrin. *Neurology.* 1999;53(9):2082-7.
23. Wild JM, Chiron C, Ahn H, et al. Visual field loss in patients with refractory partial epilepsy treated with vigabatrin: final results from an open-label, observational, multicentre study. *CNS Drugs.* 2009;23(11):965-82.
24. Vonthein R, Rauscher S, Paetzold J, et al. The normal age-corrected and reaction time-corrected isopter derived by semi-automated kinetic perimetry. *Ophthalmology.* 2007;114(6):1065-72.
25. Bengtsson B, Heijl A. False-negative responses in glaucoma perimetry: indicators of patient performance or test reliability? *Invest Ophthalmol Vis Sci.* 2000;41(8):2201-04.



26. Manning CD, Raghavan P, Schütze H. Introduction to Information Retrieval. Cambridge University Press. 2008. Cambridge, UK.
27. Bracewell RN. The Fourier Transform and its applications. McGraw Hill. 2000. New York, NY.
28. Heijl A, Lindgren A, Lindgren G. Test-retest variability in glaucomatous visual fields. *Am J Ophthalmol*. 1989;108(2):130-5.
29. Wild JM, Pacey IE, O'Neill EC, Cunliffe IA. The SITA perimetric threshold algorithms in glaucoma. *Invest Ophthalmol Vis Sci*. 1999;40(9):1998-2009.
30. Besch D, Schiefer U, Eter N, et al. Modelling the topography of absolute defects in patients exposed to the anti-epileptic drug vigabatrin and in normal subjects using automated static suprathereshold perimetry of the entire 80° visual field. *Graefes Arch Clin Exp Ophthalmol*. 2011;249(9):1333-43.
31. Westall CA, Wright T, Cortese F, Kumarappah A, Snead OC 3rd, Buncic JR. Vigabatrin retinal toxicity in children with infantile spasms: An observational cohort study. *Neurology*. 2014;83(24):2262-8.
32. Harding GFA, Wild JM, Robertson KA, et al. Separating the retinal electrophysiologic effects of vigabatrin. Treatment versus field loss. *Neurology*. 2000;55(3):347-52.
33. Origlieri C, Geddie B, Karwoski B, et al. Optical coherence tomography to monitor vigabatrin toxicity in children. *J AAPOS*. 2016;20(2):136-40.
34. Wild JM, Robson CR, Jones AL, Cunliffe IA, Smith PE Detecting vigabatrin toxicity by imaging of the retinal nerve fiber layer. *Invest Ophthalmol Vis Sci*. 2006;47(3):917-24.
35. Lawthom C, Smith PE, Wild JM. Nasal retinal nerve fiber layer attenuation: a biomarker for vigabatrin toxicity. *Ophthalmology*. 2009;116(3):565-71.

36. Clayton LM, Dévilé M, Punte T, et al. Retinal nerve fiber layer thickness in vigabatrin-exposed patients. *Ann Neurol*. 2011;69(5):845-54.
37. Clayton LM, Dévilé M, Punte T, et al. Patterns of peripapillary retinal nerve fiber layer thinning in vigabatrin-exposed individuals. *Ophthalmology*. 2012;119(10):2152-60.
38. Wild JM, Aljarudi S, Smith PEM, Knupp C. The topographical relationship between visual field loss and peripapillary retinal nerve fibre layer thinning arising from long-term exposure to vigabatrin. *CNS Drugs*. 2019;33(2):161-73.
39. Wright T, Kumarappah A, Stavropoulos A, Reginald A, Buncic JR, Westall CA. Vigabatrin toxicity in infancy is associated with retinal defect in adolescence: A prospective observational study. *Retina*. 2017;37(5):858-66.
40. Lawden MC, Eke T, Degg C, Harding GF, Wild JM. Visual field defects associated with vigabatrin therapy. *J Neurol Neurosurg Psychiatry*. 1999;67(6):716-22.
41. Frisén L, Malmgren K. Characterization of vigabatrin-associated optic atrophy. *Acta Ophthalmol Scand*. 2003;81(5):466-73.
42. Sabril Starter Kit, SHARE. Lundbeck Inc. Deerfield, IL.
43. Frisén L. Identification of functional visual field loss by automated static perimetry. *Acta Ophthalmol*. 2014;92(8):805-809.

## Figure Captions

**Fig 1** The number of individuals by anti-epileptic drug exposure, type of perimetry and visual field outcome. Equivocal indicates a normal outcome to the Central 30-2 Threshold Test; however, in the absence of the Full Field 135 Point Screening Test with the three zone age-corrected strategy, an evaluation for VAVFL was not possible. VAVFL indicates vigabatrin-associated visual field loss

**Fig 2** Stages One and Two of vigabatrin-associated visual field loss for the Full Field 135 Point Screening Test with the three zone age-corrected strategy and for the Central 30-2 Threshold Test

For the Full Field 135 Point Screening Test, ○ represents a ‘seen’ response to the initial (dimmet) stimulus luminance; × represents a ‘non-seen’ response to the initial (dimmet) stimulus luminance but a ‘seen’ response to the maximum (brightest) stimulus luminance; ■ represents a ‘non-seen’ response to the maximum (brightest) stimulus luminance

For the Central 30-2 Threshold Test, MD indicates Mean Deviation, PSD indicates Pattern Standard Deviation and SI indicates Symmetry Index. The symbols ::, ⦿, ⦿ and ■ indicate the probability of the difference between the measured value of sensitivity and the corresponding age-corrected normal value, after the general height adjustment, lying within the statistically normal range at  $p < 5\%$ ,  $p < 2\%$ ,  $p < 1\%$  and  $p < 0.5\%$ , respectively

**Fig 3** Stages Three and Four of vigabatrin-associated visual field loss for the Full Field 135 Point Screening Test with the three zone age-corrected strategy and for the Central 30-2 Threshold Test.

For the Full Field 135 Point Screening Test, ○ represents a ‘seen’ response to the initial (dimmet) stimulus luminance; × represents a ‘non-seen’ response to the initial (dimmet) stimulus luminance but a ‘seen’ response to the maximum (brightest) stimulus luminance; ■ represents a ‘non-seen’ response to the maximum (brightest) stimulus luminance

For the Central 30-2 Threshold Test, MD indicates Mean Deviation, PSD indicates Pattern Standard Deviation and SI indicates Symmetry Index. The symbols ::, ⦿, ⦿ and ■ indicate the probability of the difference between the measured value of sensitivity and the corresponding age-corrected normal value, after the general height adjustment, lying within the statistically normal range at  $p < 5\%$ ,  $p < 2\%$ ,  $p < 1\%$  and  $p < 0.5\%$ , respectively

**Fig 4** Stages Five and Six of vigabatrin-associated visual field loss for the Full Field 135 Point Screening Test with the three zone age-corrected strategy and for the Central 30-2 Threshold Test.

For the Full Field 135 Point Screening Test, ○ represents a ‘seen’ response to the initial (dimmiest) stimulus luminance; × represents a ‘non-seen’ response to the initial (dimmiest) stimulus luminance but a ‘seen’ response to the maximum (brightest) stimulus luminance; ■ represents a ‘non-seen’ response to the maximum (brightest) stimulus luminance

For the Central 30-2 Threshold Test, MD indicates Mean Deviation, PSD indicates Pattern Standard Deviation and SI indicates Symmetry Index. The symbols ::, ⌘, ⌘ and ■ indicate the probability of the difference between the measured value of sensitivity and the corresponding age-corrected normal value, after the general height adjustment, lying within the statistically normal range at  $p < 5\%$ ,  $p < 2\%$ ,  $p < 1\%$  and  $p < 0.5\%$ , respectively

**Fig 5** The Mean Deviation and Pattern Standard Deviation as a function of the ranked outcome of the Central 30-2 Threshold Test for each eye of the 99 individuals. The circles represent the Mean Deviation and the squares the Pattern Standard Deviation. The black symbols represent the outcome from the right eye and the open symbols that from the left eye

**Fig 6.** The two components of the symmetry index. The coloured area comprises the data points from one million randomly generated pairs of fields, each of which possesses differing within- and between-eye levels of field loss and represents the likelihood of the symmetry outcome occurring due to chance at  $p > 0.01$ . The black circles represent individuals exposed to vigabatrin, the vast majority of which lie above the shaded area indicating a high level of symmetry. Note the circles in the bottom left hand corner indicate an absence of symmetry since the fields are normal and those in the top right corner indicate a high degree of symmetry since the fields exhibit advanced loss. The scale on the right hand ordinate indicates the level of probability attributable to a random occurrence of symmetry: any circle lying within the white region exhibits a  $p \leq 0.01$  of symmetry occurring due to chance

**Fig 7** A case of progressive VAVFL within the central field of each eye, illustrated in terms of the Pattern Deviation probability map. The visual fields up to 82 months from onset of therapy were obtained using the Central 24-2 Threshold Test and subsequently with the Central 30-2 Threshold Test. Vigabatrin was withdrawn after 93 months of therapy. The patient remained asymptomatic. MD indicates Mean Deviation and PSD indicates Pattern Standard Deviation. The outcome of the Full Field 135 Point Screening Test at 101 months is given in Online Resource Fig 5

## Staging of vigabatrin-associated visual field loss

**Objective derivation of the morphology and staging of visual field loss associated with long-term vigabatrin therapy**

John M Wild<sup>1</sup>, Phillip E M Smith<sup>2</sup>, Carlo Knupp<sup>1</sup>

<sup>1</sup> College of Biomedical Sciences, Cardiff University, Maindy Road, Cardiff CF24 4HQ, United Kingdom

<sup>2</sup> Alan Richens Unit, Welsh Epilepsy Centre, University Hospital of Wales, Heath Park, Cardiff, CF14 4XW, United Kingdom.

**Corresponding Author:** John Wild.

**Address:** College of Biomedical Sciences, Cardiff University, Maindy Road, Cardiff CF24 4HQ, United Kingdom.

**Telephone:** ++44 29 20 87 64 87

**Fax:** ++44 29 20 87 48 59

**Email:** [wildjm@cardiff.ac.uk](mailto:wildjm@cardiff.ac.uk)

**John Wild:** ORCID: 0000-0003-3019-3889

## Abstract

**Background** The morphology and between-eye symmetry of the visual field loss associated with the anti-epileptic drug vigabatrin (VAVFL) has received little attention.

**Objective** To model the appearance, and ensuing staging, of VAVFL, derived with the European Medicines Agency approved perimetric protocol.

**Methods** A retrospective, cross-sectional, observational design identified 123 adults who had received vigabatrin for refractory seizures and who had no evidence of co-existing retino-geniculo-cortical visual pathway abnormality. Thirty-eight adults with refractory seizures and identical inclusion criteria, but **no exposure** to vigabatrin, acted as controls. For each group, the median outcome at each stimulus location in each eye (of absolute loss, relative loss or Pattern Deviation probability level, as appropriate) was derived for each successive ten pairs of fields, ranked for severity. Between-eye symmetry was quantified by an index which accounted for severity of loss and which was referenced to the likelihood of the occurrence of symmetry due to chance.

**Results** The modelled VAVFL was bilateral and highly symmetrical and was described by six stages which were all independent of the extent of vigabatrin exposure. The loss originated in the extreme temporal periphery and encroached centripetally along all meridians towards fixation. The initial appearance within the central field (Stage Two) occurred inferior-nasally. Subsequent stages exhibited increasing loss which was greater nasally than temporally. Stage Six described concentric loss extending to approximately 15° eccentricity from fixation.

**Conclusion** The model exhibited a consistent pattern of VAVFL. The staging of the loss could assist the risk:benefit analysis of vigabatrin for the treatment of epilepsy.

**Key Points**

In this study of 123 individuals treated with vigabatrin as adjunct therapy for refractory focal seizures and without evidence of other retino-geniculo-cortical visual pathway abnormality, objective modelling, based upon convolution theory used in signal-to-noise processing, identified a consistent pattern of bilateral symmetrical visual field loss obtained using a regulatory approved perimetric protocol. No modelled field loss was present for the 38 individuals with identical inclusion criteria but no exposure to vigabatrin.

Six stages of modelled vigabatrin-associated field loss (VAVFL) were evident. Originating in the extreme temporal periphery, the field loss encroached, with increasing severity, centripetally along all meridians towards fixation with greater nasal than temporal loss. Stage Six manifested as concentric loss to within approximately 15° from fixation.

The between-eye mirror symmetry of the VAVFL at all stages, quantified by a novel index, which accounted for the severity of loss, was not attributable to chance ( $p \leq 0.01$ ).

The use of either the Central 30-2 or the Central 24-2 Threshold Tests, alone, will not identify VAVFL until late Stage Two and late stage Three, respectively.

## 1.0 Introduction

The anti-epileptic drug vigabatrin has long been associated with visual field loss (VAVFL) [1-2] which manifests as a bilateral concentric peripheral defect over a continuum of severity and which, generally, also involves the central field to varying extents [3-4]. The field loss is considered to be slowly progressive [5-7] irreversible but non-progressive on withdrawal from vigabatrin [8-9] and asymptomatic until severe loss is present [3, 10-12]. The modelled prevalence of VAVFL in adults increases rapidly after the first two years (2kg cumulative dose) of exposure [13-14] and can reach 75-80% after approximately six years (5kg cumulative dose) [14]. Vigabatrin is now available generically.

The association between vigabatrin and ocular toxicity is currently being re-evaluated [15-17] on the basis that the attribution of the toxicity emanates from retrospective studies which do not include a pre-treatment baseline evaluation. The lack of evidence for the toxicity in the two prospective studies alongside the finding that approximately 25% of individuals suitable for vigabatrin exhibited clinically significant pre-existing reduced visual acuity and/ or visual field loss and/ or peripapillary retinal nerve fibre layer thinning, has led to the suggestion that abnormalities of the afferent system are a co-morbidity of severe refractory epilepsy and are unrelated to vigabatrin [15-16]. However, the limited exposures to vigabatrin and the poor retention of individuals, within these two studies [15-16], together with inappropriate visual field methodologies and analyses, have not facilitated detection of the toxicity.

Surprisingly, there has been little emphasis on the areal and depth characteristics, the between eye-symmetry, and the staging, of VAVFL. Such information is essential not only to identify the



onset and the progression of the VAVFL but also to differentiate it from other types of (concomitant) field loss.

The areal extent and the depth of any type of visual field loss is highly dependent upon the perimetric technique used for the examination. VAVFL has been investigated by a variety of methods [7, 10-11, 16, 18-23] but the characteristics of the field loss by each technique have received little attention. Unfortunately, due to the peripheral nature of the field loss, there is no one ideal technique, across the current range of commercially available perimeters, for the assessment of VAVFL. The most pragmatic approach is that approved by the European Medicines Agency (EMA) [23] namely, Three Zone (two luminance level) suprathreshold perimetry, referenced to the age-corrected threshold, of the peripheral field out to a maximum temporal extent of 90° eccentricity and standard automated perimetry of the central field (SAP).

Given the on-going re-evaluation of the association between vigabatrin and visual field loss and the wider availability of the drug, a description of the morphology and staging of the VAVFL resulting from the EMA approved protocol, and derived from individuals with long-term exposures to vigabatrin, would be of benefit to clinicians and to patients, alike. If such a description was based upon a technique which reduced the signal-to-noise ratio and was derived from individuals with no evidence of retino-geniculo-cortical visual pathway abnormality, except that of potential vigabatrin toxicity, the characteristics of the field loss attributable to vigabatrin would become manifest from backgrounds of any unexpected field loss and of any increased variability associated with the response by the patient. Such an approach would also enable the use of the visual fields from individuals with long-term exposures to vigabatrin but with no pre-treatment baseline.

The aim of the study, therefore, was threefold: to model the morphology of VAVFL obtained with the EMA approved perimetric protocol and resulting from long-term vigabatrin therapy; to quantify any between-eye symmetry; and to describe the ensuing staging of the field loss.

## 2.0 Methods

### 2.1 Case series

A retrospective case series comprising 161 adults with refractory complex partial (focal) seizures was compiled from those presenting to the Alan Richens Unit of the Welsh Epilepsy Centre, University Hospital of Wales, Cardiff, UK, and to associated clinics. All individuals had undergone part or all of the EMA approved protocol with the Humphrey Field Analyzer 750 (Carl Zeiss, Meditec, Dublin, CA) namely, the Full Field 135 Point Test with the three zone age-corrected strategy (FF135) and the Central 30-2 Threshold Test (C30-2T) with the FASTPAC strategy. None exhibited ocular or intra-cranial conditions resulting in, or likely to cause, visual field loss, as determined by ophthalmological and neurological examination including whole-brain magnetic resonance imaging. Further exclusion criteria comprised the presence of structural and/or functional ocular abnormalities likely to impede or confound the outcome of perimetry and which are described elsewhere [24]. There was no conscious selection bias. Individuals were included on the basis of sequential retrieval of cases which met the inclusion/exclusion criteria.

## 2.2 Vigabatrin exposure

Of the 161 individuals (Fig 1, top row), 123 had been exposed to vigabatrin as add-on therapy. The remaining 38 individuals had received a variety of other antiepileptic drugs and were included as age-matched controls.

## 2.3 Perimetry

Sixty-three of the 123 individuals exposed to vigabatrin and all 38 of those exposed to other antiepileptic drugs had undergone the complete EMA approved protocol (Fig 1, second row). Of the remaining 60 individuals exposed to vigabatrin, 24 had undergone the FF135, only, and 36 had undergone the C30-2T, only (Fig 1, second row) as ordered by the treating neurologist on clinical grounds. Thus, 87 of the 123 individuals exposed to vigabatrin had undertaken the FF135 and 99 individuals had undertaken the C30-2T.

The visual fields from the first occasion, after the first visit, at which reliable perimetric outcomes had been achieved were used for the individuals in each of the two groups. The reliability criteria comprised  $\leq 20\%$  incorrect responses to the fixation loss catch trials and  $\leq 15\%$  incorrect responses to the false-positive catch trials. A criterion of  $\leq 30\%$  incorrect responses was used for the false-negative catch trials although this was widened with increasing severity of the field loss [25]. All individuals had been seizure free for a minimum of 24 hours prior to the visual field examination.

## 2.4 Morphology of visual field loss

The Single Field Printouts of the FF135 and of the C30-2T were exported to a personal computer via a Humphrey Field Analyzer 820 and saved in Tagged Image File Format. The outputs from each eye for both types of perimetry were then read by custom Naïve Bayesian character recognition software [26].

The fields were separately sorted for each group, and for each type of perimetry, as right and left eye pairs, and ranked from the least to the most affected field. The severity of the given pair of fields was defined as the sum, across the two eyes, of the number of defects, weighted to account for relative or absolute loss, respectively, derived by the FF135 and/ or for the probability level of the Pattern Deviation probability map derived by the C30-2T. In the case of those undergoing both tests, the outcome from the C30-2T was more heavily weighted.

The median outcome for each group at each stimulus location in each eye for each type of perimetry was then separately calculated for each successive ten pairs of fields, ranked for severity. Such an approach was based upon convolution theory used in signal processing [27].

The procedure resulted in 54 pairs of modelled fields from the 63 individuals exposed to vigabatrin who had undergone the combined EMA protocol; 78 pairs for the FF135 and 90 for the C30-2T. Twenty-nine pairs of modelled fields resulted from the 38 individuals with no vigabatrin exposure. The outcomes for the right and left eyes, separately, were then displayed in terms of Audio Video Interleaved (AVI) movies.

## 2.5 Between-eye mirror symmetry of visual field loss

The between-eye mirror symmetry of the field was expressed in terms of a single index which comprised two components, each of which was a fraction. The index accounted for the severity of loss and was compared to the likelihood of obtaining symmetrical field loss due to chance.

The numerator and denominator of the first component constituted of, respectively, the number of mirror image stimulus locations between the two eyes which exhibited either relative or absolute loss for the FF135, or abnormality at  $p \leq 0.02$  by Pattern Deviation probability analysis for the C30-2T, and the total number of locations exhibiting abnormality across the two eyes. The numerator and denominator of the second component comprised, respectively, the total number of locations exhibiting abnormality across the two eyes and the total number of stimulus locations across the two eyes. The index was compared to that obtained from the simulation of one million randomly generated pairs of abnormal fields which manifested varying locations, areas and depths of field loss both within- and between-eyes. The concept was such that symmetry would become apparent when compared to a series of pairs of fields with varying characteristics of asymmetric field loss.

The symmetry index was separately calculated for each pair of peripheral, and each pair of central fields, for the modelled fields and for the measured fields.

## 2.6 Staging of the measured fields for those exposed to vigabatrin

The staging of the VAVFL was derived from the modelled fields. The stages were empirically selected on the basis of clinically meaningful intervals of peripheral field loss and, subsequently, of central field loss. These intervals, in turn, were based upon the magnitude of the between-examination physiological variability in the differential light sensitivity as a function of severity

of loss [28-29]. The derivation ensured continuity across stages. The staging of the modelled peripheral fields was undertaken masked to that of the modelled central fields. The outcome of the staging of the central and peripheral fields, with respect to one another, was then validated based upon the outcome of the model from the individuals who had undergone the combined protocol. The staging was then applied to each of the 87 pairs of measured fields from the FF135 and to each of the 99 measured fields from the C30-2T.

### 3.0 Results

#### 3.1 Demographic characteristics of the case series

Of the 123 individuals exposed to vigabatrin, 50 were male and 73 female. Thirty-five were receiving vigabatrin at the time of perimetry. Of the 38 age-matched controls, 17 were male and 21 female. The mean ages of the two groups, at the time of perimetry, were 40.8 years (SD 13.6) and 38.5 years (SD 10.0), respectively.

#### 3.2 Morphology of VAVFL

The rolling medians of the 78 pairs of modelled fields for the FF135 from those exposed to vigabatrin; of the 90 pairs for the C30-2T; and of the 54 pairs for the combined EMA protocol are given in video format in Online Resource Figs 1, 2 and 3, respectively.

The rolling medians of the 29 pairs of modelled fields for the combined EMA protocol from those with no exposure to vigabatrin are given in video format in Online Resource Fig 4. The modelled fields showed was no evidence of visual field loss.

### 3.3 Staging of VAVFL

The staging of the FF135 and of the C30-2T modelled fields for those exposed to vigabatrin is shown in Figs 2 to 4.

Sixteen of the 36 individuals exposed to vigabatrin, who had undergone the C30-2T, only, manifested a normal measured central field (Fig 1 third row). These 16 fields were included in the model, but, in the absence of a peripheral field assessment, the individuals could not be categorised as to the potential outcome of the toxicity and were considered as equivocal.

Of the remaining 107 individuals exposed to vigabatrin (Fig 1, fourth row), 12 exhibited normal measured fields and 95 field loss which conformed to the modelled fields and which was therefore designated as VAVFL.

The frequency, by stage, of VAVFL is shown in Fig 1 (bottom row). Fourteen of the 95 individuals with VAVFL exhibited Stage One VAVFL, 21 Stage Two, 14 Stage Three, 22 Stage Four, 10 Stage Five and 14 Stage Six.

The relationship between the stage of loss and the summary measures of the visual field, Mean Deviation (MD) and Pattern Standard Deviation, for the C30-2T measured fields is shown in Fig 5.

### 3.4 Between-Eye Mirror Symmetry of VAVFL

The two components of the symmetry index for the FF135 and for the C30-2T are shown in Online Resource Videos 1, 2 and 3; at each stage of the modelled VAVFL in Figs 2 to 4; and for each measured field of those exposed to vigabatrin in Fig 6.

### 3.5 Demographics of VAVFL

The difference in the proportion with VAVFL by gender, 39 out of 44 males and 56 out of 63 females, was not statistically significant.

The mean age, at the time of perimetry, of those with VAVFL, 41.6 years (SD 14.4), was also similar to that of those exposed to vigabatrin but with normal fields, mean 37.9 years (SD 7.9).

Those with VAVFL manifested a greater cumulative dose (mean 7.94kg; SD 4.45; range 1.1 to 20.7) and a longer duration of therapy (8.86 years; SD 3.51; range 0.66 to 16.05) at the time of perimetry than those without field loss (mean 3.36kg; SD 4.77; range 0.11 to 16.14; and mean 3.62 years; SD 3.94; range 0.33 to 11.83; respectively). The difference between means (Student's t-test for two independent samples) were 4.58 kg (95% CI 1.71-7.45;  $p < 0.001$ ) and 5.25 years (95% CI 2.98-7.53;  $p < 0.001$ ), respectively.

There was no evidence of a relationship between the stage of the VAVFL and the extent of the exposure to vigabatrin at the time of perimetry. The median exposures at the time of



perimetry which resulted in Stage One loss were 7.7kg cumulative dose (IQR 4.8, 12.1; range 1.5-13.3) and 9.4 years (IQR 6.2, 12.1; range 2.4-16.1) and which resulted in Stage 6 loss were 6.7kg cumulative dose (IQR 4.4, 10.6; range 2.8-19.0) and 7.7 years (IQR 5.3, 10.6; range 3.1-13.0).

There was also no association between the stage of VAVFL and either age, age at onset of epilepsy or age at onset of vigabatrin.

#### 4.0 Discussion

The objective rolling median and symmetry index outcomes from the FF135 and the C30-2T confirm that vigabatrin is associated with symmetrical bilateral visual field loss. The VAVFL manifests over a range of severities that can be described in six stages. The field loss originates in the extreme temporal periphery and, as the severity increases, encroaches centripetally along all meridians towards fixation resulting in bi-nasal loss and relative temporal sparing until the end stage (Stage Six) which manifests as a concentric loss to within approximately 15° from fixation. The initial appearance in the central field (Stages Two and Three) is primarily inferior-nasally. VAVFL exhibits a high degree of between-eye mirror symmetry at each stage of loss. In contrast, the modelled field for the age-matched control individuals with no exposure to vigabatrin was entirely normal.

The apparent absence of initial loss superior-nasally within the central field is most likely attributable to the greater defect depth in this region required to achieve statistical significance. The latter, in turn, arises from the wider distribution of normal values arising

from the between-individual variation in the position of the upper eyelid and from the increased variability in response associated with the steeper gradient of the superior visual field, compared to those inferior-nasally.

The fundamental strengths of this retrospective study were the robust exclusion of individuals with retino-geniculo-cortical visual pathway abnormality; the inclusion of long-term exposures to vigabatrin (median 8.8 years; IQR 5.7 to 11.3; range 0.1 to 16.1) many of which are unparalleled in the literature; and the utilisation of the two novel objective techniques which have not previously been applied to perimetry. These latter techniques enabled objective descriptions of the characteristics and between-eye mirror symmetry of VAVFL.

The modelled field at all six stages is in agreement with that derived from cross-sectional retrospective evidence with kinetic perimetry [4]. The initial manifestation of the VAVFL in this latter model was defined as a ‘non-seen’ response to the Goldmann V4e isopter at 80° temporally and/ or at 40° nasally. This definition is entirely consistent with the location and depth of the loss described as Stage One in the current study: the absolute loss designated by the FF135 is equivalent to the Goldmann V4e stimulus. The increasing encroachment into the nasal field with relative sparing of the temporal field, during Stages Two to Five, is compatible with the model out to approximately 80° temporally, derived from cross-sectional evidence, using suprathreshold perimetry equivalent to the I4e isopter [30].

The model was developed from the visual fields of adults. The reduction in amplitude of the 30Hz flicker cone electroretinogram (ERG) in infants treated with vigabatrin for infantile spasms [31] is compatible with that for adults manifesting VAVFL [32]. Similarly, the

topographical characteristics of the reduced peripapillary retinal nerve fibre layer thickness in children [33] is also compatible with that found in adults [34-38]. Those with vigabatrin-associated 30Hz flicker cone ERG abnormality in infancy subsequently manifest VAVFL, and corresponding retinal nerve fibre layer thinning, in later childhood/ early adolescence [39]. There is no reason to suggest, therefore, that the vigabatrin toxicity manifested in infancy will result in a different appearance to the VAVFL when the latter measure is obtained in later life.

The ophthalmological examinations were undertaken by any one of four ophthalmologists, depending upon the particular clinic, who were all unaware of the findings from their colleagues. All four were aware of the anti-epileptic drug history and, usually, of the visual field outcome. However, the modelling of the visual fields from those exposed to vigabatrin and from the control individuals was objective and independent of the outcome from the ophthalmological and neurological examinations.

The most common bilateral finding by fundoscopy, through a dilated pupil, was either generalised or localised arteriolar narrowing which was noted at all stages of VAVFL. However, this finding was occasionally found in the presence of a normal field and, therefore, may well be associated with vigabatrin usage. Subtle bilateral retinal nerve fibre layer changes and optic nerve head pallor were noted from Stage 4 onwards. A variety of bilateral peripheral degenerative changes were also noted including hypopigmented/ white spots and surface wrinkling. However, there was considerable variation in the fundal appearance between-individuals within a given stage. The various features are in agreement with those reported previously [3, 22, 40-41], were often subtle, and suggest that there is a

wide spectrum of potential retinopathy associated with vigabatrin toxicity. There was no evidence of such findings in the individuals with no exposure to vigabatrin. Twenty-nine individuals exposed to vigabatrin had undergone optical coherence tomography of the peripapillary retinal nerve fibre layer at the time of the examination. All three individuals who manifested a normal visual field exhibited a normal nerve fibre layer thickness. Of the three individuals who had Stage 1 loss, two exhibited a normal thickness. An abnormally thin nerve fibre layer, characteristic of vigabatrin toxicity [34-38], was present from Stage 2 onwards in 22 individuals. The one remaining individual manifested early Stage 2 VAVFL and a normal a normal nerve fibre layer thickness.

The outcome of the current study and of the earlier models [4, 28] indicates that perimetry out to 60° eccentricity [42] is inadequate to detect Stage One and also possibly early Stage Two loss. Similarly, a worsening of 3dB from the baseline Mean Deviation (MD) index for the C30-2T, which was used as a primary outcome measure for the presence of VAVFL in the prospective Phase IV study [16], will not detect VAVFL until late in Stage Two or early in Stage Three. The MD is a summary measure of the central field and does not describe the spatial appearance of visual field loss: in the current study, the median MDs for the individuals exposed to vigabatrin and for the controls was -1.75dB and -1.23dB, respectively.

It is clear from the staging of the VAVFL that perimetry should initially be undertaken with, in the case of the Humphrey Field Analyzer, the FF135. If the field is normal to the FF135 or exhibits Stage One abnormality, the C30-2T is unnecessary. If the peripheral field exhibits Stage Two abnormality to the FF135, the C30-2T should also be undertaken to reveal the full depth of the loss since the magnitude of the Pattern Deviation value required for statistical

significance at the extreme nasal locations within the central field can be less than the 8dB suprathereshold increment of the stimulus luminance of the FF135. If the FF135 indicates both peripheral and central loss, i.e., Stage Three and worse, subsequent examinations need only be undertaken with the C30-2T. The Central 24-2 Threshold Test, alone, should only be used for Stage Six. The characteristics of VAVFL are more prominent with the FASTPAC algorithm than with the SITA Standard or Fast algorithms which were designed to detect glaucomatous field loss. The peripheral manifestation of VAVFL is fortuitous since, in our experience, patients exposed to vigabatrin find suprathereshold perimetry considerably easier to perform than SAP. In cases where patients are unable to undertake SAP, assessment with the FF135, although collecting less information about relative loss, is acceptable.

The staging of VAVFL was derived from cross-sectional evidence and does not imply progressive loss. Due to the potency of the potential toxicity, most individuals had been withdrawn from vigabatrin either immediately prior to the introduction into the care regime of the visual field examination or following confirmation of the VAVFL. As such, neither the probability of progression at each stage nor an estimate of the rate of progression can be determined. However, the presumption is that, given continued vigabatrin therapy, VAVFL will progress through the various stages. The lack of an association between the stage of loss, at detection, and either the duration or the cumulative dose of vigabatrin implies that the relationship between the extent of exposure and both the onset and the severity of VAVFL varies depending upon the individual susceptibility to vigabatrin. However, the time of detection is not the time to onset of the VAVFL. The rate of any subsequent progression, therefore, remains unknown. A case of progressive loss during approximately 7.75 years of vigabatrin therapy and illustrated in terms of the Pattern Deviation probability map of the Central 24-2 Threshold Test and, subsequently, the C30-2T is shown in Fig 6. The outcome

of the corresponding FF135 at the final visit is shown in Online Resource Fig 4. Progressive loss whilst on therapy has been shown over a mean follow-up of 10.7 years; the rate of reduction in the I4e isopter of kinetic perimetry increased with increase in cumulative dose [6]. An unpublished audit of the long-term follow-up, over a maximum of eight years, of individuals in the current study indicates that the VAVFL neither improves nor deteriorates following withdrawal of vigabatrin.

The appearance of Stage One VAVFL should act as an alert signal for a re-evaluation of the management of the epilepsy. This may encompass either a change in anti-epileptic drug therapy or an increase in the frequency of neuro-ophthalmological monitoring. However, the extreme peripheral stimulus locations of the FF135 are also influenced by the facial anatomy, particularly superiorly and inferior-nasally. The outcome of the examination at such locations, and those in the extreme temporal periphery, is also dependent upon the vigilance of the patient. Indeed, normal individuals do not necessarily produce statistically normal visual fields. The FF135 commences with the examination of the central field. Once completed, the examination pauses for removal of the trial lenses prior to the examination of the peripheral field. Instructions to the patient, during this pause, to the effect that the stimuli are about to appear in the (extreme) periphery, together with an exhortation to keep the eye ‘wide open’, generally leads to a ‘seen’ response, in the normal eye, at these extreme locations. However, individuals with epilepsy frequently exhibit greater within-examination variability than normal individuals, which can lead to the impression of visual field loss, and approximately 30% of individuals with epilepsy are unable to undertake perimetry of any type [15,32]. The clinical skill lies in the identification of actual visual field loss, characteristic of the given lesion, from apparent visual field loss due to increased variability

and/ or an inability to understand the requirements of the test. In addition, VAVFL frequently co-exists with that arising from an intra-cranial lesion.

Fourteen individuals exhibited Stage Six VAVFL. Great care was taken to exclude individuals manifesting visual field loss of suspected psychogenic origin from the case series [43]. Three of these 14 individuals had undergone optical coherence tomography and exhibited severe thinning of the retinal nerve fibre layer. All 14 individuals exhibited profoundly impaired mobility and reported symptoms explainable by their VAVFL, the most common of which was bumping into individuals in crowded locations. However, some individuals with Stage 5 VAVFL also reported a negative impact on their activities of daily living.

The study utilised a retrospective cross-sectional design. However, the objective derivation of field loss based upon signal-to-noise theory, further supported by the stringent ophthalmological and neurological inclusion criteria, uniquely facilitated a retrospective design and largely obviated the need for a prospective study referenced to a pre-treatment baseline.

The variable and lengthy time to onset of VAVFL together with the opportunity to study a wide range of long-term exposures, further highlights the pragmatism of this retrospective study compared to a prospective study. Nevertheless, prospective longitudinal studies over the long-term would unequivocally rule out the presence of abnormality at baseline and would also determine the evolution of the field loss.

## 5.0 Conclusions

The appearance of VAVFL has received little attention. This absence of knowledge hinders not only the recognition of the onset, and of any progression, of the field loss but also the differentiation from other (concomitant) types of loss. The staging of the VAVFL derived from a regulatory approved perimetric protocol, and reported here, should assist clinicians and patients, alike, in the risk:benefit analysis of vigabatrin for the treatment of epilepsy.

## 6.0 Compliance with Ethical Standards

**Funding** None

**Conflicts of interest** John M Wild, Phillip EM Smith and Carlo Knupp declare that they have no conflicts of interest.

**Ethical approval** All procedures performed in studies involving human participants were in accordance with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. The Local Research and Ethics Committee ruled that approval was not required for this study. The visual field assessments were considered to be part of normal good clinical practice. The visual fields were de-identified and so written informed consent was not required.

**Data availability** The datasets generated and/or analysed during the current study are available from the corresponding author on reasonable request.



## 7.0 References

1. Eke T, Talbot JF, Lawden MC. Severe persistent visual field constriction associated with vigabatrin. *BMJ*. 1997;314(7075):180-1.
2. Maguire MJ, Hemming K, Wild JM, et al. Prevalence of visual field loss following exposure to vigabatrin therapy: a systematic review. *Epilepsia*. 2010;51(12):2423-31.
3. Wild JM, Martinez C, Reinshagen G, Harding GF. Characteristics of a unique visual field defect attributed to vigabatrin. *Epilepsia*. 1999;40(12):1784-94.
4. Malmgren K, Ben-Menachem E, Frisén L. Vigabatrin visual toxicity: evolution and dose dependence. *Epilepsia*. 2001;42(5):609-15.
5. Hardus P, Verduin W, Postma G, et al. Long term changes in the visual fields of patients with temporal lobe epilepsy using vigabatrin. *Br J Ophthalmol*. 2000;84(7):788-90.
6. Clayton LM, Stern WM, Newman WD, et al. Evolution of visual field loss over ten years in individuals taking vigabatrin. *Epilepsy Research*. 2013;105(3):262-71.
7. Nowomiejska K, Jedrych M, Brzozowska A, et al. Relationship between the area of isopters and vigabatrin dosage during two years of observation. *BMC Ophthalmol*. 2014;14:56. doi: 10.1186/1471-2415-14-56.
8. Johnson MA, Krauss GL, Miller NR, et al. Visual function loss from vigabatrin: effect of stopping the drug. *Neurology*. 2000;55(1):40-5.
9. Nousiainen I, Mäntyjärvi M, Kälviäinen R. No reversion in vigabatrin-associated visual field defects. *Neurology*. 2001;57(10):1916-7.
10. Daneshvar H, Racette L, Coupland SG, et al. Symptomatic and asymptomatic visual loss in patients taking vigabatrin. *Ophthalmology*. 1999;106(9):1792-8.

11. Kälviäinen R, Nousiainen I, Mäntyjärvi M, et al. Vigabatrin, a gabaergic drug, causes concentric visual field defects. *Neurology*. 1999;53(5):922-6.
12. Hardus P, Verduin WM, Postma G, et al. Concentric contraction of the visual field in patients with temporal lobe epilepsy and its association with the use of vigabatrin medication. *Epilepsia*. 2000;41(5):581-7.
13. European Medicines Agency. Opinion of the Committee for proprietary medicinal products pursuant to Article 12 of Council Directive 75/319/EEC as amended for vigabatrin. Annex 1. Scientific conclusions and grounds for amendment of the summaries of product characteristics presented by the EMEA. 1999, European Medicines Agency, Canary Wharf, London, United Kingdom. Available at:  
[https://www.ema.europa.eu/documents/referral/opinion-committee-proprietary-medicinal-products-pursuant-article-12-council-directive-75/319/eec-amended-vigabatrin-annexes-i-ii-iii-iv\\_en.pdf](https://www.ema.europa.eu/documents/referral/opinion-committee-proprietary-medicinal-products-pursuant-article-12-council-directive-75/319/eec-amended-vigabatrin-annexes-i-ii-iii-iv_en.pdf) Accessed 6th April 2019.
14. Wild JM, Fone DL, Aljarudi S, et al. Modelling the risk of visual field loss arising from long-term exposure to the anti-epileptic drug vigabatrin: a cross-sectional approach. *CNS Drugs*. 2013;27(10):841-9.
15. Krauss G, Faught E, Foroozan R, et al. Sabril® registry 5-year results: Characteristics of adult patients treated with vigabatrin. *Epilepsy Behav*. 2016;56(3):15-19.
16. Sergott RC, Johnson CA, Laxer KD, et al. Retinal structure and function in vigabatrin-treated adult patients with refractory complex partial seizures. *Epilepsia* 2016;57(10):1634-42.
17. Foroozan R. Vigabatrin: lessons learned from the United States experience. *J Neuro-Ophthalmol*. 2018;38(4):442-50.

18. Hardus P, Verduin WM, Engelsman M, et al. Visual field loss associated with vigabatrin: quantification and relation to dosage. *Epilepsia*. 2001;42(2):262-7.
19. Kinirons P, Cavalleri GL, O'Rourke D, et al. Vigabatrin retinopathy in an Irish cohort: lack of correlation with dose. *Epilepsia*. 2006;47(2):311-7.
20. Manuchehri K, Goodman S, Siviter L, Nightingale S. A controlled study of vigabatrin and visual abnormalities. *Br J Ophthalmol*. 2000;84(5):499-505.
21. van der Torren K, Graniewski-Wijnands HS, Polak BC. Visual field and electrophysiological abnormalities due to vigabatrin. *Doc Ophthalmol*. 2002;104(2):181-8.
22. Miller NR, Johnson MA, Paul SR, et al. Visual dysfunction in patients receiving vigabatrin. *Neurology*. 1999;53(9):2082-7.
23. Wild JM, Chiron C, Ahn H, et al. Visual field loss in patients with refractory partial epilepsy treated with vigabatrin: final results from an open-label, observational, multicentre study. *CNS Drugs*. 2009;23(11):965-82.
24. Vonthein R, Rauscher S, Paetzold J, et al. The normal age-corrected and reaction time-corrected isopter derived by semi-automated kinetic perimetry. *Ophthalmology*. 2007;114(6):1065-72.
25. Bengtsson B, Heijl A. False-negative responses in glaucoma perimetry: indicators of patient performance or test reliability? *Invest Ophthalmol Vis Sci*. 2000;41(8):2201-04.
26. Manning CD, Raghavan P, Schütze H. *Introduction to Information Retrieval*. Cambridge University Press. 2008. Cambridge, UK.
27. Bracewell RN. *The Fourier Transform and its applications*. McGraw Hill. 2000. New York, NY.

28. Heijl A, Lindgren A, Lindgren G. Test-retest variability in glaucomatous visual fields. Am J Ophthalmol. 1989;108(2):130-5.
29. Wild JM, Pacey IE, O'Neill EC, Cunliffe IA. The SITA perimetric threshold algorithms in glaucoma. Invest Ophthalmol Vis Sci. 1999;40(9):1998-2009.
30. Besch D, Schiefer U, Eter N, et al. Modelling the topography of absolute defects in patients exposed to the anti-epileptic drug vigabatrin and in normal subjects using automated static suprathreshold perimetry of the entire 80° visual field. Graefes Arch Clin Exp Ophthalmol. 2011;249(9):1333-43.
31. Westall CA, Wright T, Cortese F, Kumarappah A, Snead OC 3rd, Buncic JR. Vigabatrin retinal toxicity in children with infantile spasms: An observational cohort study. Neurology. 2014;83(24):2262-8.
32. Harding GFA, Wild JM, Robertson KA, et al. Separating the retinal electrophysiologic effects of vigabatrin. Treatment versus field loss. Neurology. 2000;55(3):347-52.
33. Origlieri C, Geddie B, Karwoski B, et al. Optical coherence tomography to monitor vigabatrin toxicity in children. J AAPOS. 2016;20(2):136-40.
34. Wild JM, Robson CR, Jones AL, Cunliffe IA, Smith PE Detecting vigabatrin toxicity by imaging of the retinal nerve fiber layer. Invest Ophthalmol Vis Sci. 2006;47(3):917-24.
35. Lawthom C, Smith PE, Wild JM. Nasal retinal nerve fiber layer attenuation: a biomarker for vigabatrin toxicity. Ophthalmology. 2009;116(3):565-71.
36. Clayton LM, Dévilé M, Punte T, et al. Retinal nerve fiber layer thickness in vigabatrin-exposed patients. Ann Neurol. 2011;69(5):845-54.
37. Clayton LM, Dévilé M, Punte T, et al. Patterns of peripapillary retinal nerve fiber layer thinning in vigabatrin-exposed individuals. Ophthalmology. 2012;119(10):2152-60.

38. Wild JM, Aljarudi S, Smith PEM, Knupp C. The topographical relationship between visual field loss and peripapillary retinal nerve fibre layer thinning arising from long-term exposure to vigabatrin. *CNS Drugs*. 2019;33(2):161-73.
39. Wright T, Kumarappah A, Stavropoulos A, Reginald A, Buncic JR, Westall CA. Vigabatrin toxicity in infancy is associated with retinal defect in adolescence: A prospective observational study. *Retina*. 2017;37(5):858-66.
40. Lawden MC, Eke T, Degg C, Harding GF, Wild JM. Visual field defects associated with vigabatrin therapy. *J Neurol Neurosurg Psychiatry*. 1999;67(6):716-22.
41. Frisén L, Malmgren K. Characterization of vigabatrin-associated optic atrophy. *Acta Ophthalmol Scand*. 2003;81(5):466-73.
42. Sabril Starter Kit, SHARE. Lundbeck Inc. Deerfield, IL.
43. Frisén L. Identification of functional visual field loss by automated static perimetry. *Acta Ophthalmol*. 2014;92(8):805-809.

## Figure Captions

**Fig 1** The number of individuals by anti-epileptic drug exposure, type of perimetry and visual field outcome. Equivocal indicates a normal outcome to the Central 30-2 Threshold Test; however, in the absence of the Full Field 135 Point Screening Test with the three zone age-corrected strategy, an evaluation for VAVFL was not possible. VAVFL indicates vigabatrin-associated visual field loss

**Fig 2** Stages One and Two of vigabatrin-associated visual field loss for the Full Field 135 Point Screening Test with the three zone age-corrected strategy and for the Central 30-2 Threshold Test

For the Full Field 135 Point Screening Test, ○ represents a 'seen' response to the initial (dimmet) stimulus luminance; × represents a 'non-seen' response to the initial (dimmet) stimulus luminance but a 'seen' response to the maximum (brightest) stimulus luminance; ■ represents a 'non-seen' response to the maximum (brightest) stimulus luminance

For the Central 30-2 Threshold Test, MD indicates Mean Deviation, PSD indicates Pattern Standard Deviation and SI indicates Symmetry Index. The symbols □, ▨, ▩ and ■ indicate the probability of the difference between the measured value of sensitivity and the corresponding age-corrected normal value, after the general height adjustment, lying within the statistically normal range at  $p < 5\%$ ,  $p < 2\%$ ,  $p < 1\%$  and  $p < 0.5\%$ , respectively

**Fig 3** Stages Three and Four of vigabatrin-associated visual field loss for the Full Field 135 Point Screening Test with the three zone age-corrected strategy and for the Central 30-2 Threshold Test.

For the Full Field 135 Point Screening Test, ○ represents a 'seen' response to the initial (dimmet) stimulus luminance; × represents a 'non-seen' response to the initial (dimmet) stimulus luminance but a 'seen' response to the maximum (brightest) stimulus luminance; ■ represents a 'non-seen' response to the maximum (brightest) stimulus luminance

For the Central 30-2 Threshold Test, MD indicates Mean Deviation, PSD indicates Pattern Standard Deviation and SI indicates Symmetry Index. The symbols □, ▨, ▩ and ■ indicate the probability of the difference between the measured value of sensitivity and the corresponding age-corrected normal value, after the general height adjustment, lying within the statistically normal range at  $p < 5\%$ ,  $p < 2\%$ ,  $p < 1\%$  and  $p < 0.5\%$ , respectively

**Fig 4** Stages Five and Six of vigabatrin-associated visual field loss for the Full Field 135 Point Screening Test with the three zone age-corrected strategy and for the Central 30-2 Threshold Test.

For the Full Field 135 Point Screening Test, ○ represents a 'seen' response to the initial (dimmiest) stimulus luminance; × represents a 'non-seen' response to the initial (dimmiest) stimulus luminance but a 'seen' response to the maximum (brightest) stimulus luminance; ■ represents a 'non-seen' response to the maximum (brightest) stimulus luminance

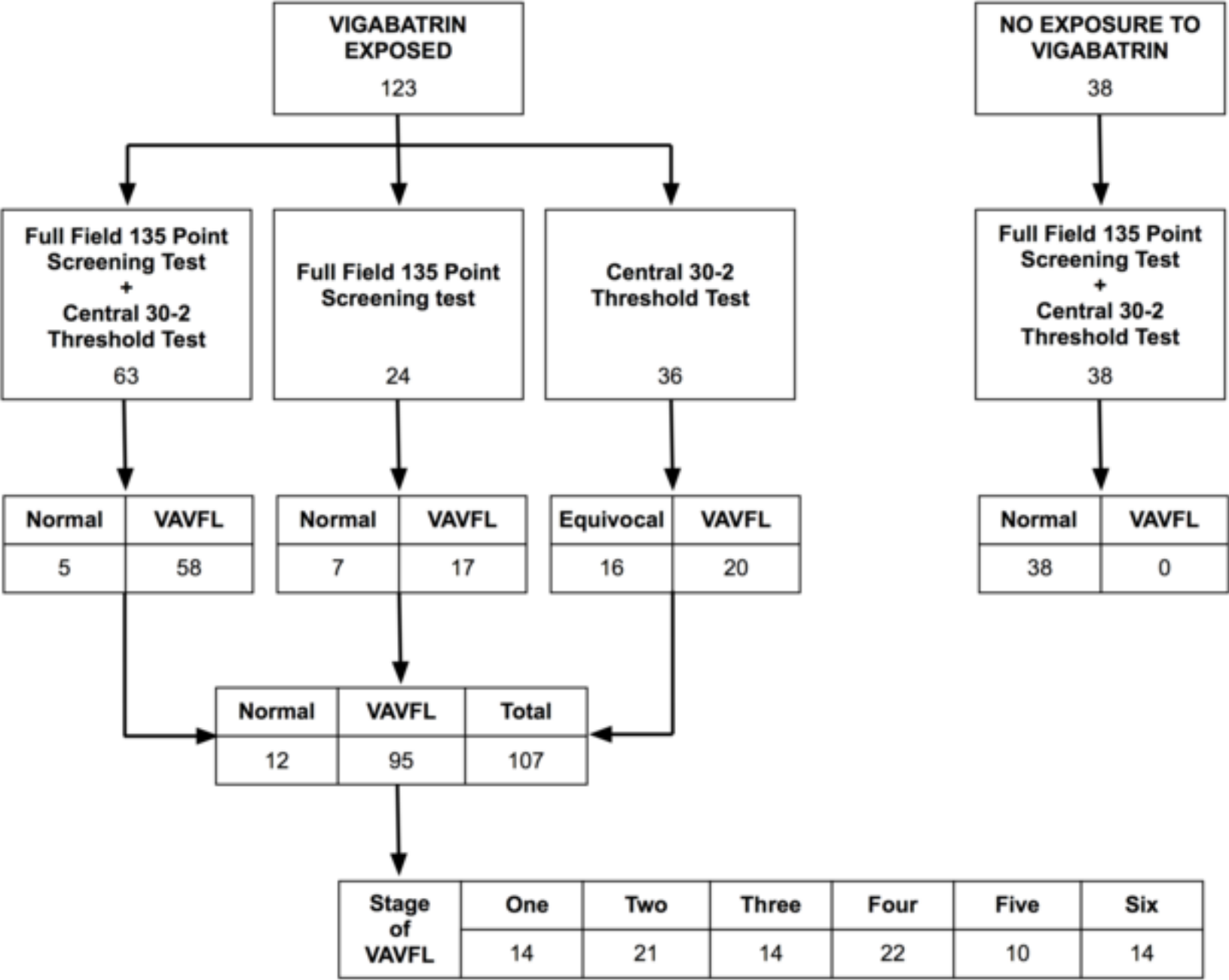
For the Central 30-2 Threshold Test, MD indicates Mean Deviation, PSD indicates Pattern Standard Deviation and SI indicates Symmetry Index. The symbols ::, ⌘, ⌘ and ■ indicate the probability of the difference between the measured value of sensitivity and the corresponding age-corrected normal value, after the general height adjustment, lying within the statistically normal range at  $p < 5\%$ ,  $p < 2\%$ ,  $p < 1\%$  and  $p < 0.5\%$ , respectively

**Fig 5** The Mean Deviation and Pattern Standard Deviation as a function of the ranked outcome of the Central 30-2 Threshold Test for each eye of the 99 individuals. The circles represent the Mean Deviation and the squares the Pattern Standard Deviation. The black symbols represent the outcome from the right eye and the open symbols that from the left eye

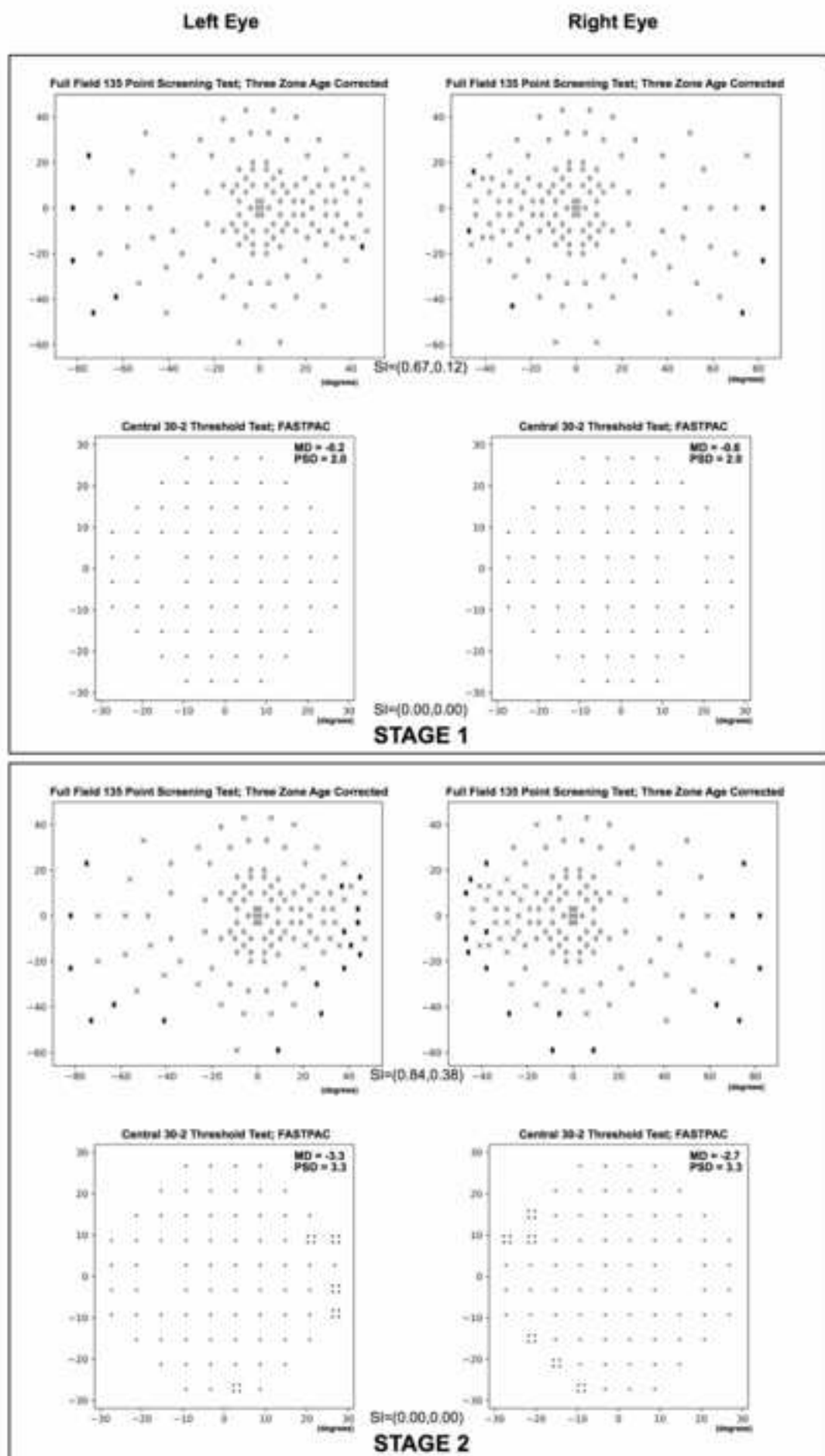
**Fig 6.** The two components of the symmetry index. The coloured area comprises the data points from one million randomly generated pairs of fields, each of which possesses differing within- and between-eye levels of field loss and represents the likelihood of the symmetry outcome occurring due to chance at  $p > 0.01$ . The black circles represent individuals exposed to vigabatrin, the vast majority of which lie above the shaded area indicating a high level of symmetry. Note the circles in the bottom left hand corner indicate an absence of symmetry since the fields are normal and those in the top right corner indicate a high degree of symmetry since the fields exhibit advanced loss. The scale on the right hand ordinate indicates the level of probability attributable to a random occurrence of symmetry: any circle lying within the white region exhibits a  $p \leq 0.01$  of symmetry occurring due to chance

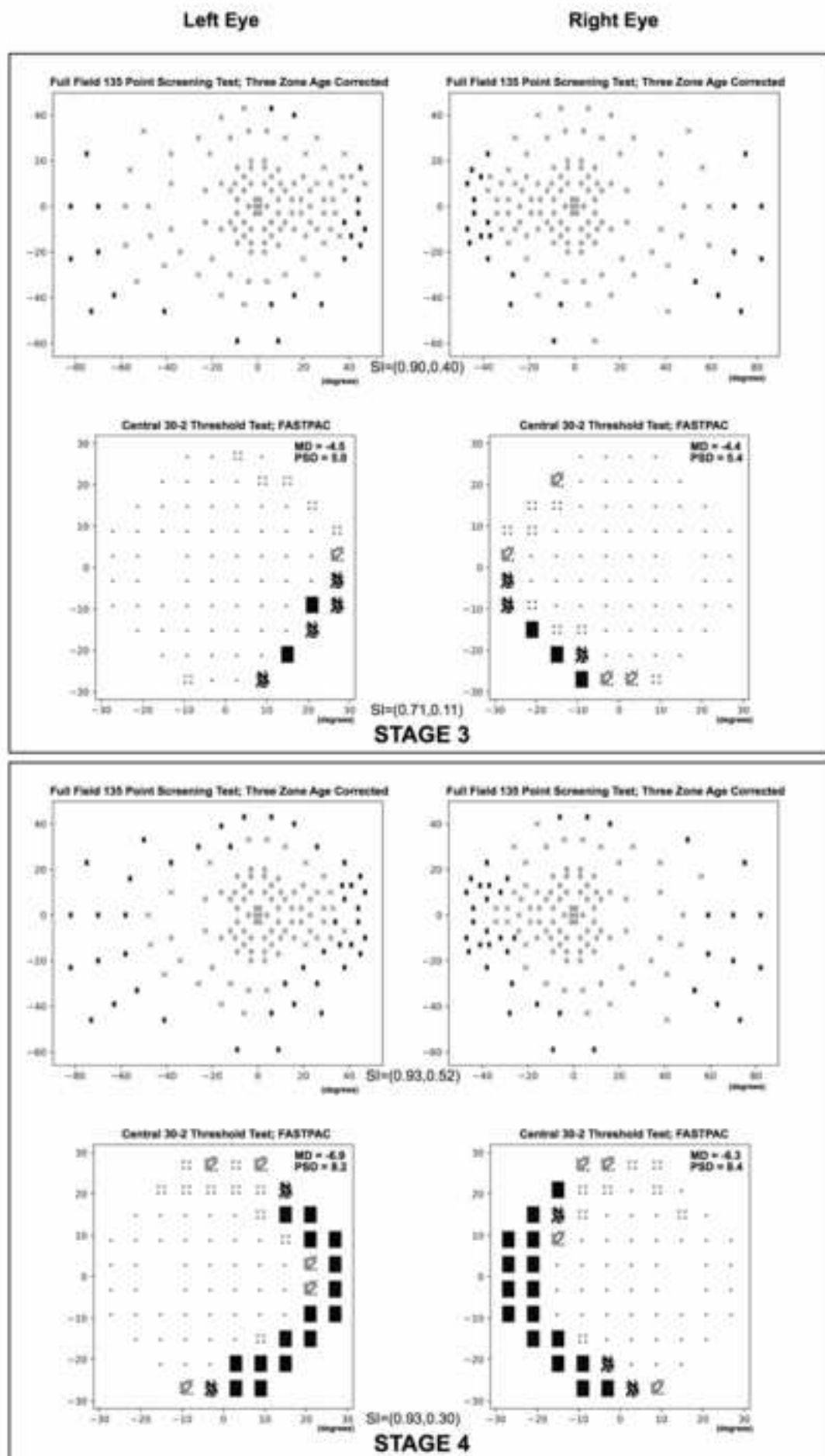
**Fig 7** A case of progressive VAVFL within the central field of each eye, illustrated in terms of the Pattern Deviation probability map. The visual fields up to 82 months from onset of therapy were obtained using the Central 24-2 Threshold Test and subsequently with the Central 30-2 Threshold Test. Vigabatrin was withdrawn after 93 months of therapy. The patient remained asymptomatic. MD indicates Mean Deviation and PSD indicates Pattern Standard Deviation. The outcome of the Full Field 135 Point Screening Test at 101 months is given in Online Resource Fig 5

Figure 1









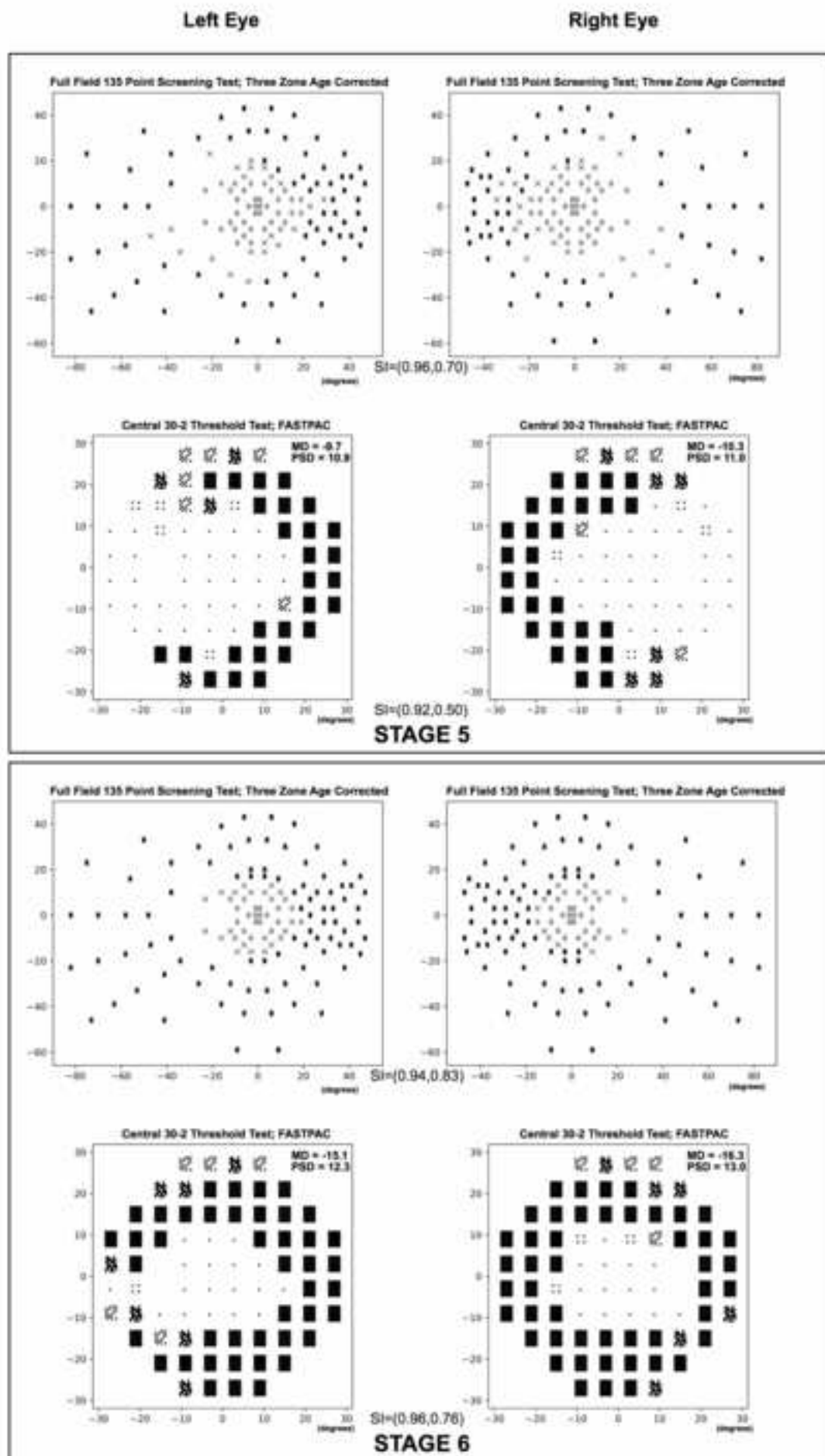
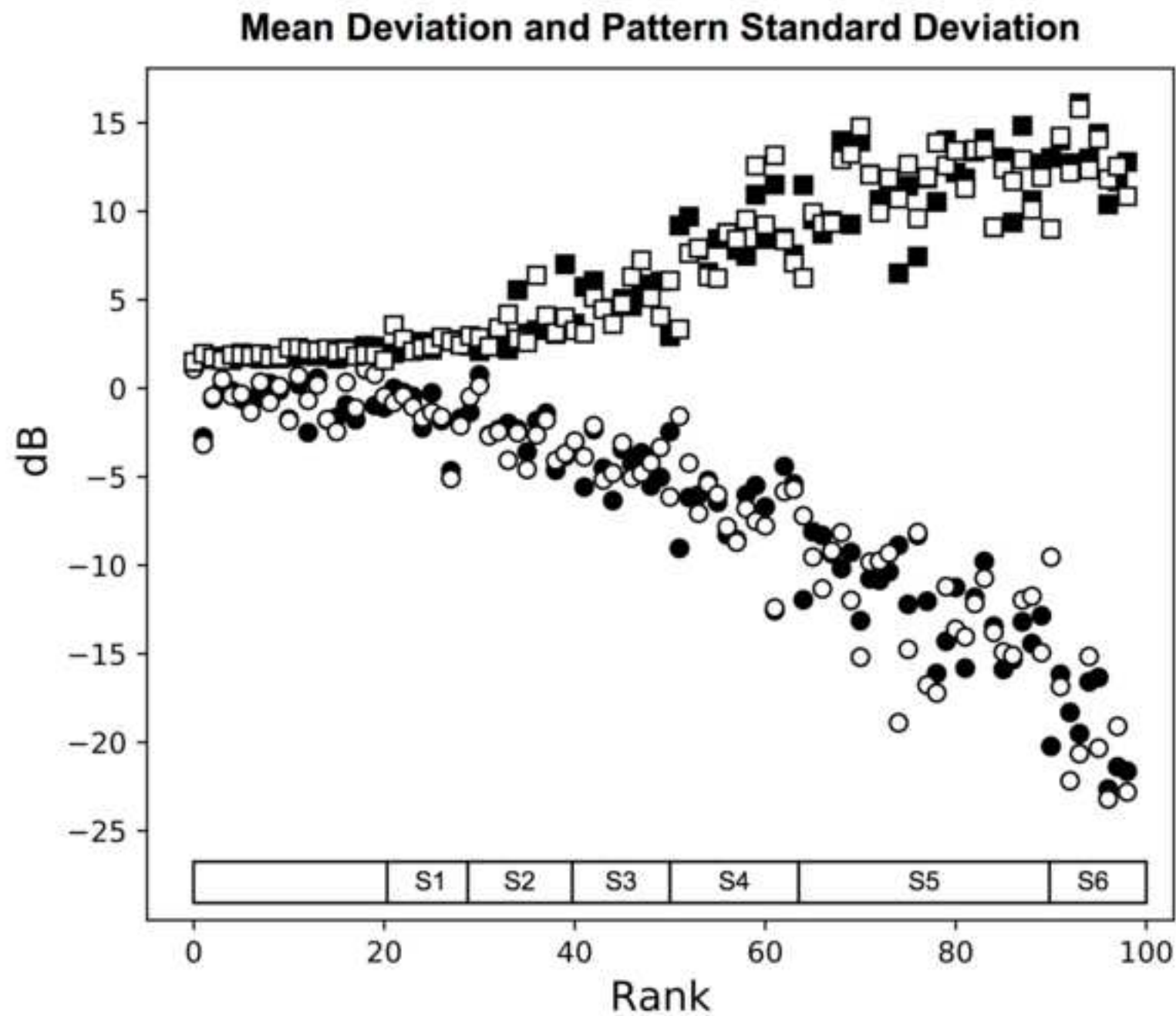
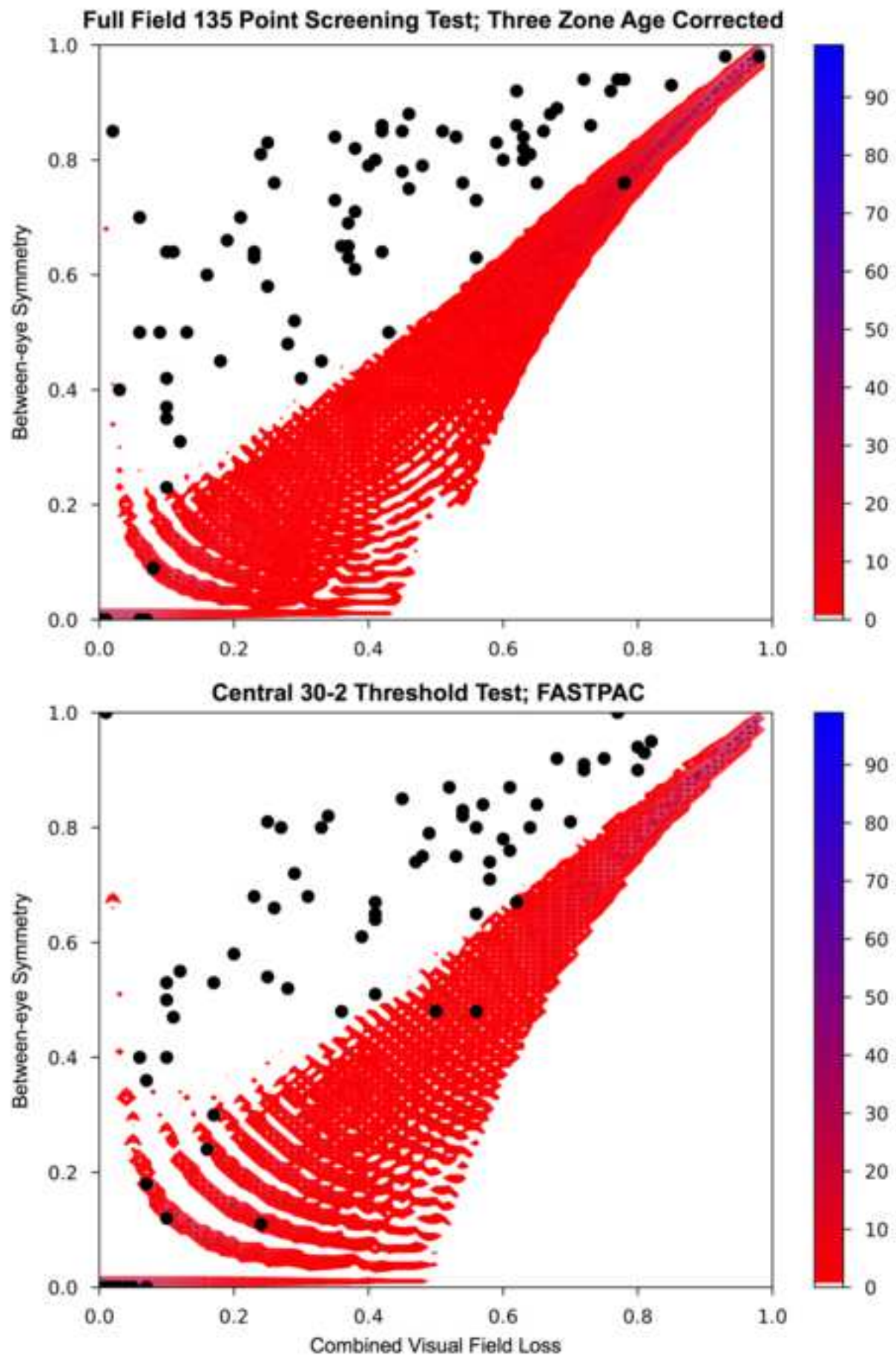
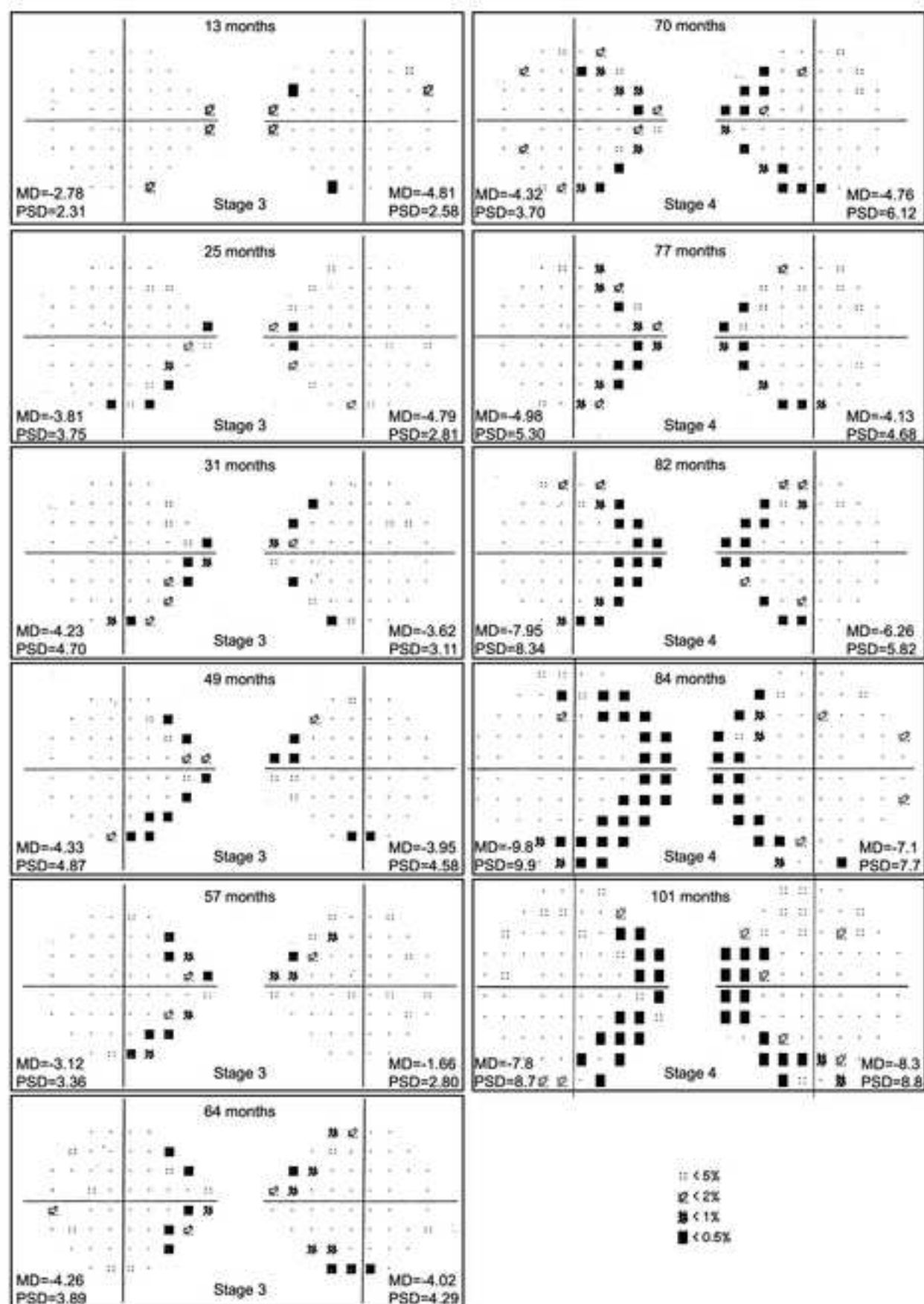


Figure 5









# **Objective derivation of the morphology and staging of visual field loss associated with long-term vigabatrin therapy**

## **CNS Drugs**

John M Wild<sup>1</sup>, Phillip E M Smith<sup>2</sup>, Carlo Knupp<sup>1</sup>

<sup>1</sup> College of Biomedical Sciences, Cardiff University, Maindy Road, Cardiff CF24 4HQ, United Kingdom

<sup>2</sup> Alan Richens Unit, Welsh Epilepsy Centre, University Hospital of Wales, Heath Park, Cardiff, CF14 4XW, United Kingdom.

**Corresponding Author:** John Wild.

**Email:** [wildjm@cardiff.ac.uk](mailto:wildjm@cardiff.ac.uk)

**Online Resource Fig 1** Video of the rolling median of the 78 modelled fields from the 87 individuals who had been exposed to vigabatrin and who had undertaken the Full Field 135 Point Test with the three zone age-corrected strategy

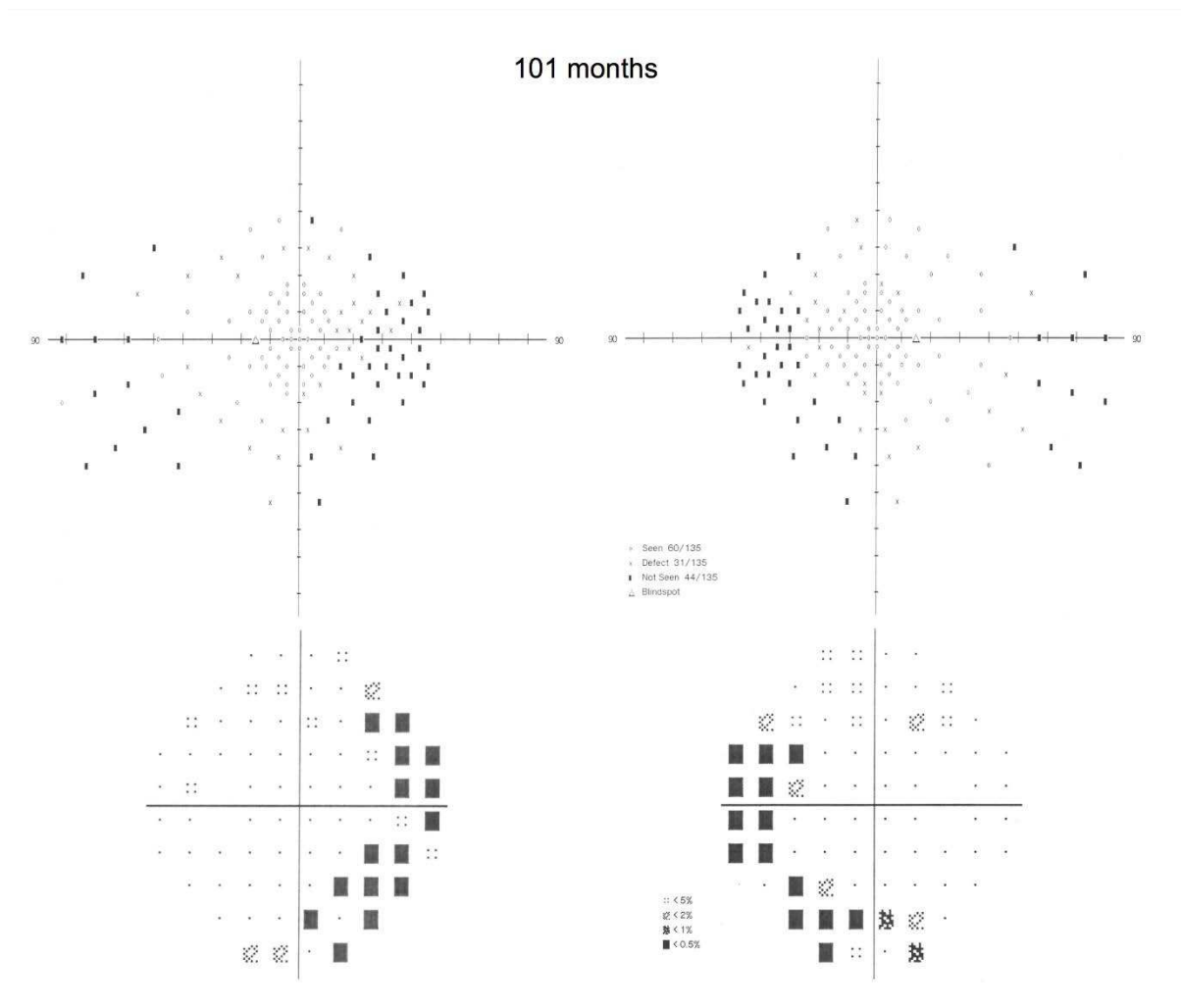
**Online Resource Fig 2** Video of the rolling median of the 90 modelled fields from the 99 individuals who had been exposed to vigabatrin and who had undertaken the Central 30-2 Threshold Test

**Online Resource Fig 3** Video of the rolling median of the 54 modelled fields from the 63 individuals who had been exposed to vigabatrin and who had undertaken the combined Full Field 135 Point Test with the three zone age-corrected strategy and the Central 30-2 Threshold Test

**Online Resource Fig 4** Video of the rolling median of the 29 modelled fields from the 38 individuals with no exposure to vigabatrin and who had undertaken the combined Full Field 135 Point Test with the three zone age-corrected strategy and the Central 30-2 Threshold Test

All video files are available here: <https://doi.org/10.6084/m9.figshare.7981592>





**Online Resource Fig 5** The outcome of the Full Field 135 Point Screening Test at 101 months for the case shown in Fig 5 together with that of the Central 30-2 Threshold Test

## AUTHOR DECLARATION FORM

At submission, **EVERY AUTHOR** listed in the manuscript must **READ** and **COMPLETE** the following statements on:  
(A) Authorship Responsibility, (B) Authorship Criteria, (C) Authorship Contribution, (D) Funding Disclosures,  
(E) Contributor Disclosures/Acknowledgments, and (F) Conflicts of Interest Disclosures.

It is important that you return this form as early as possible in the publication process. **EVERY AUTHOR MUST COMPLETE AN INDIVIDUAL COPY OF THE FORM, AND EVERY SECTION OF THE FORM MUST BE COMPLETED.**  
We will **NOT** consider your manuscript for publication until every author has completed the form and returned it to us.

Your name (please print): Carlo Knpp \_\_\_\_\_ E-mail: knuppc@cardiff.ac.uk \_\_\_\_\_

Journal name: CNS Drugs \_\_\_\_\_ Corresponding author: John Wild \_\_\_\_\_

Manuscript title: Objective derivation of the morphology and staging of visual field loss associated with long-term vigabatrin therapy

### A. AUTHORSHIP RESPONSIBILITY

☒ I certify that **ALL** of the following statements are correct (**PLEASE CHECK THE BOX**).

- The manuscript represents valid work; neither this manuscript nor one with substantially similar content under my authorship has been published or is being considered for publication elsewhere (except as described in the manuscript submission); and copies of any closely related manuscripts are enclosed in the manuscript submission; **AND**
- For manuscripts with more than one author, I agree to allow the corresponding author to serve as the primary correspondent with the editorial office and to review and sign off on the final proofs prior to publication; or, if I am the only author, I will be the corresponding author and agree to serve in the roles described above.
- For manuscripts that are a report of a study, I confirm that this work is an accurate representation of the trial results.

### B. AUTHORSHIP CRITERIA

To fulfil all of the criteria for authorship, every author of the manuscript must have made substantial contributions to **ALL** of the following aspects of the work:

- Conception and planning of the work that led to the manuscript or acquisition, analysis and interpretation of the data, or both; **AND**
- Drafting and/or critical revision of the manuscript for important intellectual content; **AND**
- Approval of the final submitted version of the manuscript.

☒ I certify that I fulfill **ALL** of the above criteria for authorship (**PLEASE CHECK THE BOX**).

### C. AUTHORSHIP CONTRIBUTION

I certify that I have participated sufficiently in the work to take public responsibility for (**PLEASE CHECK 1 OF THE 2 BOXES BELOW**):

- ☐ Part of the content of the manuscript; **OR**  
☒ The entire content of the manuscript.

### D. FUNDING DISCLOSURES

**PLEASE CHECK 1 OF THE 2 BOXES BELOW:**

- ☒ I certify that no funding has been received for the conduct of this study and/or preparation of this manuscript; **OR**  
☐ I certify that all financial and material support for the conduct of this study and/or preparation of this manuscript is clearly described in the Compliance with Ethical Standards section of the manuscript.

Some funding organizations require that authors of manuscripts reporting research deposit those manuscripts with an approved public repository.

☐ Please check here if you have received such funding.

### E. CONTRIBUTOR DISCLOSURES

All persons who have made substantial contributions to the work reported in the manuscript (e.g. data collection, data analysis, or writing or editing assistance) but who do not fulfill the authorship criteria **MUST** be named with their specific contributions in the Acknowledgments section of the manuscript. Groups of persons who have contributed may be listed under a heading such as 'Clinical investigators' and their function described. Because readers may infer their endorsement of the manuscript, all persons named in the Acknowledgments section **MUST** give the authors their written permission to be named in the manuscript.

- ☒ I certify that all persons who have made substantial contributions to this manuscript but who do not fulfill the authorship criteria are listed with their specific contributions in the Acknowledgments section in the manuscript, and that all persons named in the Acknowledgments section have given me written permission to be named in the manuscript.

## F. CONFLICT OF INTEREST DISCLOSURES

A conflict of interest exists when professional judgment concerning a primary interest (such as patients' welfare or the validity of research) may be influenced by a secondary interest (such as financial gain or personal rivalry). A conflict of interest may arise for authors when they have a financial interest that may influence – probably without their knowing – their interpretation of their results or those of others. We believe that to make the best decision on how to deal with a manuscript we should know about any such conflict of interest that the authors may have. We are not aiming to eradicate conflicts of interests – they are almost inevitable. We will not reject manuscripts simply because the authors have a conflict of interest, but we will publish a declaration in the manuscript as to whether or not the authors have conflicts of interests.

All authors **MUST** complete the following checklist:

<b>Category of potential conflict of interest</b>	If you have had any of the listed relationships with an entity that has a financial interest in the subject matter discussed in this manuscript, please check the appropriate "Yes" box below and provide details. If you do not have a listed relationship, please check the appropriate "No" box. When completing this section, please take into account the last 36 months through to the foreseeable future.		
	<b>No (✓)</b>	<b>Yes (✓)</b>	<b>Details</b>
Employment	✓		
Grant received/grants pending	✓		
Consulting fees or honorarium	✓		
Support for travel to meetings for the study, manuscript preparation or other purposes	✓		
Fees for participation in review activities such as data monitoring boards, etc	✓		
Payment for writing or reviewing the manuscript	✓		
Provision of writing assistance, medicines, equipment or administrative support	✓		
Payment for lectures including service on speakers bureaus	✓		
Stock/stock options	✓		
Expert testimony	✓		
Patents (planned, pending or issued)	✓		
Royalties	✓		
Other (err on the side of full disclosure)	✓		

Every author **MUST** complete option 1 or option 2 as appropriate below. If you answered "Yes" to any of the questions relating to financial conflicts of interests in the table above (or if you wish to disclose a non-financial conflict of interest), you **MUST** write a suitable statement in the box below and include this statement in the Compliance with Ethical Standards section of the manuscript.

☒ I have no conflicts of interest to declare; **OR**

☐ The following statement regarding conflicts of interest and financial support for conduct of this study and/or preparation of this manuscript is to be published in the Compliance with Ethical Standards section of the manuscript:

**Declaration:** I certify that I have fully read and fully understood this form, and that the information that I have presented here is accurate and complete to the best of my knowledge.

Your name (please print): Carlo Knupp\_\_\_\_\_

Signature (please **HAND-WRITE**) :  \_\_\_\_\_

Date: 14<sup>th</sup> February 2019\_\_\_\_\_



## AUTHOR DECLARATION FORM

At submission, **EVERY AUTHOR** listed in the manuscript must **READ** and **COMPLETE** the following statements on:  
(A) Authorship Responsibility, (B) Authorship Criteria, (C) Authorship Contribution, (D) Funding Disclosures,  
(E) Contributor Disclosures/Acknowledgments, and (F) Conflicts of Interest Disclosures.

It is important that you return this form as early as possible in the publication process. **EVERY AUTHOR MUST COMPLETE AN INDIVIDUAL COPY OF THE FORM, AND EVERY SECTION OF THE FORM MUST BE COMPLETED.**  
We will **NOT** consider your manuscript for publication until every author has completed the form and returned it to us.

Your name (please print): John Wild \_\_\_\_\_ E-mail: wildjm@cardiff.ac.uk \_\_\_\_\_

Journal name: CNS Drugs \_\_\_\_\_ Corresponding author: John Wild \_\_\_\_\_

Manuscript title: Objective derivation of the morphology and staging of visual field loss associated with long-term vigabatrin therapy

### A. AUTHORSHIP RESPONSIBILITY

☒ I certify that **ALL** of the following statements are correct (**PLEASE CHECK THE BOX**).

- The manuscript represents valid work; neither this manuscript nor one with substantially similar content under my authorship has been published or is being considered for publication elsewhere (except as described in the manuscript submission); and copies of any closely related manuscripts are enclosed in the manuscript submission; **AND**
- For manuscripts with more than one author, I agree to allow the corresponding author to serve as the primary correspondent with the editorial office and to review and sign off on the final proofs prior to publication; or, if I am the only author, I will be the corresponding author and agree to serve in the roles described above.
- For manuscripts that are a report of a study, I confirm that this work is an accurate representation of the trial results.

### B. AUTHORSHIP CRITERIA

To fulfil all of the criteria for authorship, every author of the manuscript must have made substantial contributions to **ALL** of the following aspects of the work:

- Conception and planning of the work that led to the manuscript or acquisition, analysis and interpretation of the data, or both; **AND**
- Drafting and/or critical revision of the manuscript for important intellectual content; **AND**
- Approval of the final submitted version of the manuscript.

☒ I certify that I fulfill **ALL** of the above criteria for authorship (**PLEASE CHECK THE BOX**).

### C. AUTHORSHIP CONTRIBUTION

I certify that I have participated sufficiently in the work to take public responsibility for (**PLEASE CHECK 1 OF THE 2 BOXES BELOW**):

- ☐ Part of the content of the manuscript; **OR**  
☒ The entire content of the manuscript.

### D. FUNDING DISCLOSURES

**PLEASE CHECK 1 OF THE 2 BOXES BELOW:**

- ☒ I certify that no funding has been received for the conduct of this study and/or preparation of this manuscript; **OR**  
☐ I certify that all financial and material support for the conduct of this study and/or preparation of this manuscript is clearly described in the Compliance with Ethical Standards section of the manuscript.

Some funding organizations require that authors of manuscripts reporting research deposit those manuscripts with an approved public repository.

☐ Please check here if you have received such funding.

### E. CONTRIBUTOR DISCLOSURES

All persons who have made substantial contributions to the work reported in the manuscript (e.g. data collection, data analysis, or writing or editing assistance) but who do not fulfill the authorship criteria **MUST** be named with their specific contributions in the Acknowledgments section of the manuscript. Groups of persons who have contributed may be listed under a heading such as 'Clinical investigators' and their function described. Because readers may infer their endorsement of the manuscript, all persons named in the Acknowledgments section **MUST** give the authors their written permission to be named in the manuscript.

- ☒ I certify that all persons who have made substantial contributions to this manuscript but who do not fulfill the authorship criteria are listed with their specific contributions in the Acknowledgments section in the manuscript, and that all persons named in the Acknowledgments section have given me written permission to be named in the manuscript.

## F. CONFLICT OF INTEREST DISCLOSURES

A conflict of interest exists when professional judgment concerning a primary interest (such as patients' welfare or the validity of research) may be influenced by a secondary interest (such as financial gain or personal rivalry). A conflict of interest may arise for authors when they have a financial interest that may influence – probably without their knowing – their interpretation of their results or those of others. We believe that to make the best decision on how to deal with a manuscript we should know about any such conflict of interest that the authors may have. We are not aiming to eradicate conflicts of interests – they are almost inevitable. We will not reject manuscripts simply because the authors have a conflict of interest, but we will publish a declaration in the manuscript as to whether or not the authors have conflicts of interests.

All authors **MUST** complete the following checklist:

<b>Category of potential conflict of interest</b>	If you have had any of the listed relationships with an entity that has a financial interest in the subject matter discussed in this manuscript, please check the appropriate "Yes" box below and provide details. If you do not have a listed relationship, please check the appropriate "No" box. When completing this section, please take into account the last 36 months through to the foreseeable future.		
	<b>No (✓)</b>	<b>Yes (✓)</b>	<b>Details</b>
Employment	✓		
Grant received/grants pending	✓		
Consulting fees or honorarium	✓		
Support for travel to meetings for the study, manuscript preparation or other purposes	✓		
Fees for participation in review activities such as data monitoring boards, etc	✓		
Payment for writing or reviewing the manuscript	✓		
Provision of writing assistance, medicines, equipment or administrative support	✓		
Payment for lectures including service on speakers bureaus	✓		
Stock/stock options	✓		
Expert testimony	✓		
Patents (planned, pending or issued)	✓		
Royalties	✓		
Other (err on the side of full disclosure)	✓		

Every author **MUST** complete option 1 or option 2 as appropriate below. If you answered "Yes" to any of the questions relating to financial conflicts of interests in the table above (or if you wish to disclose a non-financial conflict of interest), you **MUST** write a suitable statement in the box below and include this statement in the Compliance with Ethical Standards section of the manuscript.

☒ I have no conflicts of interest to declare; **OR**

☐ The following statement regarding conflicts of interest and financial support for conduct of this study and/or preparation of this manuscript is to be published in the Compliance with Ethical Standards section of the manuscript:

**Declaration:** I certify that I have fully read and fully understood this form, and that the information that I have presented here is accurate and complete to the best of my knowledge.

Your name (please print): John Wild\_\_\_\_\_



Signature (please **HAND-WRITE**): \_\_\_\_\_

Date: 14<sup>th</sup> February 2019\_\_\_\_\_

## AUTHOR DECLARATION FORM

At submission, **EVERY AUTHOR** listed in the manuscript must **READ** and **COMPLETE** the following statements on:  
(A) Authorship Responsibility, (B) Authorship Criteria, (C) Authorship Contribution, (D) Funding Disclosures,  
(E) Contributor Disclosures/Acknowledgments, and (F) Conflicts of Interest Disclosures.

It is important that you return this form as early as possible in the publication process. **EVERY AUTHOR MUST COMPLETE AN INDIVIDUAL COPY OF THE FORM, AND EVERY SECTION OF THE FORM MUST BE COMPLETED.**  
We will **NOT** consider your manuscript for publication until every author has completed the form and returned it to us.

Your name (please print): Phillip Smith\_\_\_\_\_ E-mail: smithpe@cardiff.ac.uk\_\_\_\_\_

Journal name: CNS Drugs\_\_\_\_\_ Corresponding author: John Wild\_\_\_\_\_

Manuscript title: Objective derivation of the morphology and staging of visual field loss associated with long-term vigabatrin therapy

### A. AUTHORSHIP RESPONSIBILITY

☒ I certify that **ALL** of the following statements are correct (**PLEASE CHECK THE BOX**).

- The manuscript represents valid work; neither this manuscript nor one with substantially similar content under my authorship has been published or is being considered for publication elsewhere (except as described in the manuscript submission); and copies of any closely related manuscripts are enclosed in the manuscript submission; **AND**
- For manuscripts with more than one author, I agree to allow the corresponding author to serve as the primary correspondent with the editorial office and to review and sign off on the final proofs prior to publication; or, if I am the only author, I will be the corresponding author and agree to serve in the roles described above.
- For manuscripts that are a report of a study, I confirm that this work is an accurate representation of the trial results.

### B. AUTHORSHIP CRITERIA

To fulfil all of the criteria for authorship, every author of the manuscript must have made substantial contributions to **ALL** of the following aspects of the work:

- Conception and planning of the work that led to the manuscript or acquisition, analysis and interpretation of the data, or both; **AND**
- Drafting and/or critical revision of the manuscript for important intellectual content; **AND**
- Approval of the final submitted version of the manuscript.

☒ I certify that I fulfill **ALL** of the above criteria for authorship (**PLEASE CHECK THE BOX**).

### C. AUTHORSHIP CONTRIBUTION

I certify that I have participated sufficiently in the work to take public responsibility for (**PLEASE CHECK 1 OF THE 2 BOXES BELOW**):

- ☐ Part of the content of the manuscript; **OR**  
☒ The entire content of the manuscript.

### D. FUNDING DISCLOSURES

**PLEASE CHECK 1 OF THE 2 BOXES BELOW:**

- ☒ I certify that no funding has been received for the conduct of this study and/or preparation of this manuscript; **OR**  
☐ I certify that all financial and material support for the conduct of this study and/or preparation of this manuscript is clearly described in the Compliance with Ethical Standards section of the manuscript.

Some funding organizations require that authors of manuscripts reporting research deposit those manuscripts with an approved public repository.

☐ Please check here if you have received such funding.

### E. CONTRIBUTOR DISCLOSURES

All persons who have made substantial contributions to the work reported in the manuscript (e.g. data collection, data analysis, or writing or editing assistance) but who do not fulfill the authorship criteria **MUST** be named with their specific contributions in the Acknowledgments section of the manuscript. Groups of persons who have contributed may be listed under a heading such as 'Clinical investigators' and their function described. Because readers may infer their endorsement of the manuscript, all persons named in the Acknowledgments section **MUST** give the authors their written permission to be named in the manuscript.

- ☒ I certify that all persons who have made substantial contributions to this manuscript but who do not fulfill the authorship criteria are listed with their specific contributions in the Acknowledgments section in the manuscript, and that all persons named in the Acknowledgments section have given me written permission to be named in the manuscript.



## F. CONFLICT OF INTEREST DISCLOSURES

A conflict of interest exists when professional judgment concerning a primary interest (such as patients' welfare or the validity of research) may be influenced by a secondary interest (such as financial gain or personal rivalry). A conflict of interest may arise for authors when they have a financial interest that may influence – probably without their knowing – their interpretation of their results or those of others. We believe that to make the best decision on how to deal with a manuscript we should know about any such conflict of interest that the authors may have. We are not aiming to eradicate conflicts of interests – they are almost inevitable. We will not reject manuscripts simply because the authors have a conflict of interest, but we will publish a declaration in the manuscript as to whether or not the authors have conflicts of interests.

All authors **MUST** complete the following checklist:

<b>Category of potential conflict of interest</b>	If you have had any of the listed relationships with an entity that has a financial interest in the subject matter discussed in this manuscript, please check the appropriate "Yes" box below and provide details. If you do not have a listed relationship, please check the appropriate "No" box. When completing this section, please take into account the last 36 months through to the foreseeable future.		
	<b>No (✓)</b>	<b>Yes (✓)</b>	<b>Details</b>
Employment	✓		
Grant received/grants pending	✓		
Consulting fees or honorarium	✓		
Support for travel to meetings for the study, manuscript preparation or other purposes	✓		
Fees for participation in review activities such as data monitoring boards, etc	✓		
Payment for writing or reviewing the manuscript	✓		
Provision of writing assistance, medicines, equipment or administrative support	✓		
Payment for lectures including service on speakers bureaus	✓		
Stock/stock options	✓		
Expert testimony	✓		
Patents (planned, pending or issued)	✓		
Royalties	✓		
Other (err on the side of full disclosure)	✓		


Every author **MUST** complete option 1 or option 2 as appropriate below. If you answered "Yes" to any of the questions relating to financial conflicts of interests in the table above (or if you wish to disclose a non-financial conflict of interest), you **MUST** write a suitable statement in the box below and include this statement in the Compliance with Ethical Standards section of the manuscript.

☒ I have no conflicts of interest to declare; **OR**

☐ The following statement regarding conflicts of interest and financial support for conduct of this study and/or preparation of this manuscript is to be published in the Compliance with Ethical Standards section of the manuscript:

**Declaration:** I certify that I have fully read and fully understood this form, and that the information that I have presented here is accurate and complete to the best of my knowledge.

Your name (please print): Phil Smith\_\_\_\_\_



Signature (please **HAND-WRITE**) : \_\_\_\_\_

Date: 14<sup>th</sup> February 2019\_\_\_\_\_




[Click here to access/download](#)

**Video**

[Online Resource \\_Video\\_1\\_Wild et al.avi](#)





[Click here to access/download](#)

**Video**

Online Resource \_Video\_2\_Wild et al.avi





[Click here to access/download](#)

**Video**

[Online Resource \\_Video\\_3\\_Wild et al.avi](#)





[Click here to access/download](#)

**Video**

[Online Resource \\_Video\\_4.\\_Wild et al.avi](#)

